Table 1

RECOMBINATION PROTEIN RECR [Bacillus subtilis] dehydratase, D-Ser [Escherichia coli] (Z99112) uridylate kinase [Bacillus subtilis]	85 23 49 21 82 51	3.00E-16 5.70E-16 4.80E-41	202 206 436	182 125 504	g585801 g223359 g2634023	508-369 589-328 370-1	Bt1G23 Bt1G25 Bt1G26	Bt1Gc22 Bt1Gc27 Bt1Gc29	15 · 16 17	
	50 99	1.60E-15	195	191	g586863	351-130	Bt1G22	Bt1Gc22	15	
	72 53	4.60E-59	606	584	g1710383	1-469	Bt1G21	Bt1Gc20	14	
	87 67	1.00E-61	631	756	g3123268	1-501	Bt1G20	Bt1Gc21	13	
stearothermophilus (Z99120) similar to hypothetical proteins [Bacillus subtilis]	53 44	3.60E-36	390	417	g2635675	496-1	Bt1G18	Bt1Gc19	12	
	76 22	2.60E-33	363	354	g2558482	545-279	Bt1G17	Bt1Gc18	11	
PHOSPHORYLASE) (PNP) [Bacillus stearothermophilus]										
PURINE NUCLEOSIDE PHOSPHORYLASE (INOSINE	70 31	1.70E-22	261	312	g2494052	260-1	Bt1G16	Bt1Gc18	11	
	32 73	1.60E-24	280	230	g2127819	615-1	Bt1G15	Bt1Gc16	10	
	35 22	0.00022	97	87	g1107529	1-296	Bt1G14	Bt1Gc10	. 9	
		9.30E-09	143	140	g2632778	1-296	Bt1G13	Bt1Gc10	9	
				·	(
_	65 31	6.70E-19	227	211	g2635805	524-329	Bt1G10	Bt1Gc12	∞	
(AF064522) major autolysin [Moraxella sp. CK-1]	31 49	2.70E-09	140	58	g4835715	1-461	Bt1G9	Bt1Gc6	7	
[Bacillus subtilis]					•					
	63 4	6.40E-12	175	173	g2117582	144-1	Bt1G6	Bt1Gc3	0	
subtilis]										
CARBOXYPEPTIDASE) (CPASE) (PBP5) [Bacillus					•					
		i	ç	Ç	0	į		(,	
LIGASE) (ALARS) [Bacillus subtilis]	\$\$ 3 <i>3</i>	1 20E-35	3. 85	365	o585034	426-1	Brigs	Rt1Gc4		
stearott ALAN	38 15	8.90E-15	200	152	g3122886	387-1	Bt1G4	Bt1Gc8695	4	
	34 16	4.30E-11	165	112	g1944414	1-388	Bt1G3	Bt1Gc8693	w	
		•) !		, •	
	34 33	2.30E-09	145	150	g2497382	1-422	Bt1G2	Bt1Gc8696	2	
PREPROTEIN TRANSLOCASE SECA SUBUNIT	42 26	2.10E-19	238	196	g3122850	371-1	Bt1G1	Bt1Gc8697	1	
NCBI gi description	Ident Cyrg	Prob I	Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	8 B	
		,	المريدات	88t			٠		SEQ	

Table 1

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56 Bt	55 Bt	54 B	53 B	52 B	52 Bi	51 B	50 Bt	49 Bt	48 B		47 B		‡		43 B1	42 Bi		40 Bı	39 5		38 B	SEQ Co
Bt1Gc100	Bt1Gc101	Bt1Gc99	Bt1Gc98	Bt1Gc97	Bt1Gc97	Bt1Gc95	Bt1Gc92	Bt1Gc93	Bt1Gc91		Briggen	Bt1Gc87	#02011G	10.84	Bt1Gc82	Bt1Gc79	Bt1Gc75	Bt1Gc77	9/30/19		Bt1Gc73	Contig Id
Bt1G79	Bt1G78	Bt1G77	Bt1G76	Bt1G75	Bt1G74	Bt1G73	Bt1G72	Bt1G70	Bt1G69		Religion	Bt1G65	#00 IId	B+1064	Bt1G63	Bt1G62	Bt1G61	Bt1G60	BUGSY		Bt1G58	Gene Id
396-1	566-93	1-297	572-35	1-527	1-527	35-611	1-987	1-441	1-377	1	1-200	1-299	1474	1_404	223-446	59-505	417-1	1-463	1-066	3	365-1	Position
g2612883	g2621742	g2145962	g2828523	g4226092	g4490571	g3123166	g1805464	g1673402	g2495457	Br. 0.000	9990000	g121563	810/39/2	~1075077	g2635847	g3582242	g1545847	g2635913	8323/183		g1176701	NCBI gi
234	102	292	511	104	89	.282	550	189	479	į	156	274	331	351	248	67	239	481	439	3	425	aat_nap
281	159	250	487	178	146	306	533	230	495	į	178	291	0/0	370	267	133	263	462	482		156	BlastP Score
1.30E-24	1.10E-11	2.40E-21	1.90E-46	3.90E-13	2.60E-10	2.90E-27	2.50E-51	1.60E-18	2.70E-47	1.000	1 00E-57	1.10E-25	2.10E-33	2 10E-22	1.20E-22	6.10E-09	2.40E-22	8.40E-44	0.4UE-40		3.20E-19	BlastP- Prob
48	27	57	57	30 .	30	34	40	33	77	d	40 6	56	ŧ	7	63	33	40	60	2	3	71	% Ident (
90	%	38	38	4	43	28	61	30	32	į	45	27	3	ž	16	87	32	50	9	;	55	% Cvrg
(AF015825) unknown [Bacillus subtilis]	[Mycobacterium leprae] (AE000846) epoxidase [Methanobacterium	PERMEASE) [Bacillus subtilis] probable transport protein abc1 - Mycobacterium leprae	[Caelorilabulis elegans] GABA PERMEASE (4-AMINO BUTYRATE TRANSPORT CARRIER) (GAMA-AMINOBUTYRATE	(AF125448) contains similarity to methyltransferases	INTERGENIC REGION [Bacillus subtilis] (AJ010302) mg protoporphyrin methyltransferase (Rhodobacter sphaeroides)	HYPOTHETICAL 25.7 KD PROTEIN IN GERAC-FHUC	(D50453) homologue of copper export protein PcoD of E.	(Z82044) hypothetical 54.4 kd protein [Bacillus subtilis]	HYPOTHETICAL 41.5 KD PROTEIN IN AMHX-AMYE INTERGENIC REGION [Bacillus subtilis]	influenzae]	(MJ010132) hypothetical protein [Bachilus cereus]	SPORE PROTEASE [Bacillus subtilis]	perfringens]	[Bacillus subtilis]	[Lactococcus lactis] (Z99121) similar to ABC transporter (amino acid permease)	(AE001272) L. lactis predicted coding region ORF00033	(U50396) WbpA [Pseudomonas aeruginosa]	(Z99121) similar to hypothetical proteins [Bacillus subtilis]	(Arounds) 3 14aa long nypothetical protein [ryrococcus horikoshii]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 23.6 KD PROTEIN IN QCRC-DAPB	NCBI gi description

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	_	_			g419952	1-731	Bt1G100	Bt1Gc117	73
MBL PROTEIN [Bacillus cereus]		1.70E-09	144		g417314	591-498	Bt1G99	Bt1Gc135	72
(AJ010138) hypothetical protein [Bacillus cereus]	96 52	6.30E-07	114 6	116	g4584138	68-1	Bt1G98	Bt1Gc135	72
[Mycobacterium leprae] (U67061) pullulanase [Bacteroides thetaiotaomicron]	51 34	1.70E-52	544 1	607	g1561763	1-687	Bt1G97	Bt1Gc136	. 71
5] (AL049478) putative DNA polymerase III alpha chain	50 24	6.90E-72	734 6	679 .	g4539136	843-1	Bt1G96	Bt1Gc131	70
tuberculosisj HYPOTHETICAL PROTEIN UL125 [human herpesvirus	100 99	4.60E-52 1	540 4	533	g137016	141-447	Bt1G95	Bt1Gc123	69
[Streptomyces coelicolor] (AL021287) hypothetical protein Rv2994 [Mycobacterium	23 41	0.00019	107	125	g2791591	549-1	Bt1G94	Bt1Gc124	68
(AL022374) putative ATP-dependent DNA helicase	53 11	1.70E-20	252 1	234	g3036880	245-1	Bt1G92	Bt1Gc119	67
(D90900) hypothetical protein [Synechocystis sp.]	37 86	1.40E-20	243 1		g1651880	553-146	Bt1G91	Bt1Gc118	66
DNA binding protein HU [Bacillus caldolyticus]	83 .44	1.20E-12	168 1		g227370	1-121	Bt1G90	Bt1Gc118	66
(AB005149) orf2 [Exiguobacterium acetylicum]	36 99	3.80E-16	201 3		g2641974	592-168	Bt1G89	Bt1Gc116	65
(AF047044) putative transposase [Anabaena PCC7120]	22 36	4.30E-09	140 4	93	g3005554	663-318	Bt1G88	Bt1Gc111	2
IN HEMY-GLTT INTERGENIC REGION (ORFA)									•
INTERGENIC REGION [Bacillus subtilis] HYPOTHETICAL TRANSCRIPTIONAL REGULATOR	22 81	1.40E-05	103 1	89	g418449	62-523	Bt1G87	Bt1Gc115	63
HYPOTHETICAL 37.6 KD PROTEIN IN GNTR-HTPG	33 . 36	3.00E-10	151 3	111	g1176967	425-1	Bt1G86	Bt1Gc110	62
(TRANSCRIPTASE ALPHA CHAIN) (RNA POLYMERASE ALPHA SUBUNIT) [Bacillus subtilis]			٠						
Clostridium magnum [Clostridium magnum] DNA-DIRECTED RNA POLYMERASE ALPHA CHAIN	88 81	3.90E-110	1088 3.9	1123 1	g133395	55-820	Bt1G85	Bt1Gc112	61
[Bacillus stearothermophilus] TPP-dependent acetoin dehydrogenase alpha chain -	51 . 23	1.90E-21	251 1	233 :	g2148094	1-234	Bt1G84	Bt1Gc114	60
IIABC COMPONENT) (PHOSPHOTRANSFERASE ENZYME II, ABC COMPONENT) (EII-GLC / EIII-GLC)									
thermoautotrophicum] PTS SYSTEM, GLUCOSE-SPECIFIC IIABC COMBONENT (EILABC GLO) (GLUCOSE BE	60 50	2.30E-48	505 2	484	g1172714	1-488	Bt1G82	Bt1Gc105	59
[Bacıllus subtilis] (AE000804) ferritin like protein (RsgA) [Methanobacterium	46 85	2.20E-27	307 2	345	g2621200	114-553	Bt1G81	Bt1Gc104	58
(Z99112) signal recognition particle (docking protein)	75 53	5.50E-63	643 5	604	g2633967	524-1	Bt1G80	Bt1Gc103	57
NCBI gi description	nt Cvrg	BlastP- % Prob Ident		nap Score Score	NCBI gi	Position	Gene Id	Contig Id	NO E
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93 Bt1Gc179	91 Bt1Gc176 92 Bt1Gc177	89 Bt1Gc174 90 Bt1Gc171	88 Bt1Gc167		85 Bt1Gc165	84 Bt1Gc163	83 Bt1Gc152	82 Bt1Gc159	81 Bt1Gc157	80 Bt1Gc155			78 Bt1Gc153	78 Bt1Gc153	77 Bt1Gc145	76 Bt1Gc139		75 Bt1Gc134	74 Bt1Gc137	ID Contig Id
9 Bt1G124	6 Bt1G122 7 Bt1G123	4 Bt1G120 1 Bt1G121	7 Bt1G118		5 Bt1G115	3 Bt1G114	2 Bt1G113	9 Bt1G112	7 Bt1G111	5 BtlGl10			3 Bt1G108	3 Bt1G107	5 Bt1G105	9 Bt1G104	4 Bt1G103	4 Bt1G102	7 Bt1G101	d Gene Id
1-468	509-1 1-361	1-290 1-737	595-130	48-636	1-349	317-416	590-1	1-210	186-1	732-1		1-595	808-557	1-392	1-635	120-525	274-501	1-242	290-1	Position
g3183519	g121562 g3183562	g2226166 g3023410	g1001794	g4584494	g1731085	g4584153	g1903038	g132987	g1361340	g2224770	(g3915055	g4321580	g3123299	g730600	g586809	g1730988	g118716	g4468689	NCBI gi
672	671 527	338 478	105	186	255	120	597	313	92	961		596	79	480	785	398	262	242	331	nap Score
655	681 543	357 457	181	256	257	129	620	331	100	898		581	114	495	808	425	262	188	336	Score
3.00E-64	5.20E-67 2.20E-52	3.40E-32 2.80E-43	5.00E-14	5.70E-22	4.40E-22	1.60E-08	1.50E-60	6.40E-30	0.00012	5.30E-90		2.10E-56	3.60E-06	2.70E-47	1.80E-80	7.00E-40	1.30E-22	3.20E-14	1.50E-29	Prob
85	76 89	£ 65	26	34	4	79	60	89 .	49	76		56	57	69	67	57	66	67	68	Ident Cyrg
36	46	17 56	92 5		74	97	_	39	18	55			7	28	45	97	24	22	13	• •
[Bacillus subtilis] GID PROTEIN [Bacillus subtilis]	AMINO ACID UPTAKE CARRIER) [Bacillus subtilis] SPORE PROTEASE [Bacillus megaterium] SUCCINYL-COA SYNTHETASE BETA CHAIN (SCS-	(Y14080) hypothetical protein [Bacillus subtilis] BRANCHED-CHAIN AMINO ACID TRANSPORT	(D64005) hypothetical protein [Synechocystis sp.]	(AL049587) hypothetical protein [Streptomyces coelicolor]	HYPOTHETICAL 17.9 KD PROTEIN IN GLNQ-ANSR	(AJ010112) spore germination protein, GerPF [Bacillus	(Z93102) hypothetical 30.6 kd protein [Bacillus subtilis]	aureus] 50S RIBOSOMAL PROTEIN L5 []	Ilm protein - Staphylococcus aureus [Staphylococcus	(Z97025) product similar to E. coli PhoH protein [Bacillus subtilis]	HOMOLOG [Bacillus subtilis]	EXTRAGENIC SUPPRESSOR PROTEIN SUHB	CSGA INTERGENIC REGION (ORF1) [Bacillus subtilis] (AF050114) alginate Ivase [Pseudomonas sp. W7]	HYPOTHETICAL TRANSPORT PROTEIN IN NDHF-	AMINO-ACID PERMEASE ROCC [Bacillus subtilis]	HYPOTHETICAL 15.6 KD PROTEIN IN PURA-DNAC	PROBABLE METHYLTRANSFERASE [Bacillus subtilis]	[Streptomyces coelicolor] DNAJ PROTEIN [Bacillus subtilis]	(fragment) [Latrodectus tredecimguttatus] (AL035591) ABC excision nuclease subunit C	NCBI gi description

Table 1

131 132	130	128 129	127	126	126	125	124	123	122	121	120	;	119	118	117	116	115	NO BEQ
Bt1Gc253 Bt1Gc254	Bt1Gc252	Bt1Gc231 Bt1Gc251	Bt1Gc244	Bt1Gc239	Bt1Gc239	Bt1Gc240	Bt1Gc238	Bt1Gc237	Bt1Gc233	Bt1Gc232	Bt1Gc230		Bt1Gc223	Bt1Gc225	Bt1Gc226	Bt1Gc221	Bt1Gc219	Contig Id
Bt1G165 Bt1G166	Bt1G164	Bt1G162 Bt1G163	Bt1G161	Bt1G160	Bt1G159	Bt1G158	Bt1G157	Bt1G156	Bt1G154	Bt1G153	Bt1G152		Brigisi	Bt1G150	Bt1G149	Bt1G148	Bt1G147	Gene Id
1-507 1-564	115-1	1-669 352-1	1-754	655-740	1-574	1-453	1-564	1-746	1-682	381-1	562-1		386-158	301-473	500-1	409-1	1-138	Position
g1684650 g2635912	g585351	g3025461 g1731061	g1881228	g2127278	g98515	g484466	g2226166	g1731017	g2632602	g1945676	g2632528	61010001	01648861	g1176995	g2815343	g732321	g2495571	NCBI gi
400 543	142	128 470	379	129	896	72	514	584	33	171	531	. }))	128	303	373	161	aat_ nap Score
) 436 3 554	2 158	8 139 0 478	9 437	145	5 912		4 473	\$ 617	335	194	1 463		141	3 156	3 319	363	175	BlastP. Score
4.80E-41 1.50E-53	1.40E-11	1.20E-08 1.70E-45	3.70E-41	3.30E-10	1.70E-91	4.10E-09	5.70E-45	3.20E-60	2.40E-30	2.10E-15	6.60E-44		6 40E-09	3.00E-11	1.20E-28	2.60E-33	2.70E-13	BlastP- Prob
1 50 3 52	1 77	8 23 5 77	1 34	0 97	1 97		5 55) 50		33	4 53		ر م	1 52	35	61	62	% Ident
36 65	17	49 81	55	14	68	12	32	35	100	74	. 40	i	_ %	23	46	. 76	16	% Cvrg
subtilis] (Z82987) unknown [Bacillus subtilis] (Z99121) similar to transcriptional regulator (LysR family)	INTERGENIC REGION [Bacillus subtilis] CYTIDYLATE KINASE (CK) (CYTIDINE MONOPHOSPHATE KINASE) (CMP KINASE) [Bacillus		[Bacillus thuringiensis alesti] (AB001488) SIMILAR TO ENZYMES WHICH ACT VIA AN ATP-DEPENDENT COVALENT BINDING OF AMP TO THEIR SUBSTRATE. [Bacillus subtilis]	flagellin chain A - Bacillus thuringiensis (subsp. alesti)	flagellin B chain - Bacillus thuringiensis [Bacillus	glucan endo-1,3-beta-D-glucosidase (EC 3.2.1.39) precursor	(Y14080) hypothetical protein [Bacillus subtilis]	HYPOTHETICAL 80.1 KD PROTEIN IN SODA-COMGA INTERGENIC REGION [Bacillus subtilis]	(Z99105) ycgJ [Bacillus subtilis]	(Z94043) hypothetical protein [Bacillus subtilis]	(Z99105) similar to sodium/proton-dependent alanine	carboxyvinyltransferase [Bacillus subtilis]	PROTEIN IN IDH 3'REGION [Bacillus subtilis]	[Streptomyces coelicolor] HYPOTHETICAL ABC TRANSPORTER ATP-BINDING	(AL021529) putative NTP pyrophosphohydrolase	influenzae Rd] HYPOTHETICAL 22.0 KD PROTEIN IN FLIT-SECA	(THREONINE DEAMINASE) [Escherichia coli] HYPOTHETICAL PROTEIN HI0105 [Haemophilus	NCBI gi description

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NCBI gi nap Score Score g1711354 196 268	nap Score Prob Score 196 268 7.60E-23
	268 7.60E-23 731 2.60E-72 140 1.20E-08
	2 5 8 6 2
MCBI gi description Ident Cvrg [Bacillus subtilis] 47 26 PROBABLE D-SERINE DEHYDRATASE (D-SERINE DEAMINASE) [Bacillus subtilis] 64 100 (Z73234) YneB [Bacillus subtilis] 28 34 (AE000833) 5-methylcytosine-specific restriction enzyme McrB related protein [Methanobacterium thermoautotrophicum] 73 45 (AF008220) YtfP [Bacillus subtilis] 74 58 (Z99117) similar to protease [Bacillus subtilis] 75 63 42 (Z99105) similar to sodium/proton-dependent alanine	0005700575

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169	169	168	167	167		166	100	;			165	•		<u>1</u> 2		163		162	161	161	160		159	158	157	156	155	154	153		NO E)
Bt1Gc312 Bt1Gc313	Bt1Gc312	Bt1Gc311	Bt1Gc310	Bt1Gc310	. Z.	Bt1Gc308	Bt10c308				Bt1Gc305			Bt1Gc303	•	Bt1Gc302		Bt1Gc301	Bt1Gc300	Bt1Gc300	Bt1Gc296		Bt1Gc299	Bt1Gc298	Bt1Gc294	Bt1Gc297	Bt1Gc295	Bt1Gc293	Bt1Gc285		Contig Id	
Bt1G211 Bt1G212	Bt1G210	Bt1G209	Bt1G208	Bt1G207		Bt1G206	C070119				Bt1G204			Bt1G203		Bt1G202		Bt1G201	Bt1G200	Bt1G199	Bt1G198		Bt1G197	Bt1G196	Bt1G195	Bt1G194	Bt1G193	Bt1G192	Bt1G191		Gene Id	
402-525 1-809	1-324	1-726	1-1128	1-216		471-655	1-430				125-670			451-1		479-694		1-659	462-553	308-1	16-552		530-799	427-1	957-297	910-1	254-595	755-190	620-120		Position	
g225559 g113811	g4104605	g2632987	g4240001	g2632024	•	g3025068	g1/30889				g3123238			g732327		g2984723		g1684651	g2340009	g2340010	g2635183		g481591	g2635763	g3602952	g3821797	g2073397	g2407933	g4894313		NCBI gi	
1 87 1120	229	891	.168	159	٠.	100	260				589			492		300		432	93	211	297		94	513	233	815	223	209	215		nap Score	
203 1070	263	898	271	167		128	2/0				628			491		325		314	111	224	290		129	518	294	835	195	220	282		BlastP Score	
1.40E-15 3.10E-108	1.00E-22	5.30E-90	4.70E-23	1.50E-12	٠	4.30E-08	1.90E-23				2.20E-61			7.10E-47	٠	2.80E-29		4.00E-28	1.00E-05	1.40E-18	1.40E-25		1.60E-08	9.80E-50	5.30E-26	2.50E-83	1.60E-15	3.70E-18	1.00E-24		BlastP- Prob	
98 72	46	70	34	49		37	39	3			75			60		89		37	65	43	38		28	74	31	52	44	28	37		% J	
9 52	47	53	36	52		25	ŏ	3			48		•	48		99		56	7	59	50	•	47	31	94	58	98	61	99		cvrg	•
sakei] ORF IS231C [Bacillus thuringiensis] ALPHA-AMYLASE PRECURSOR (1,4-ALPHA-D-	methyltransferase [Bacillus subtilis] (AF036967) putative response regulator [Lactobacillus	(Z99107) alternate gene name: yerS; similar to RNA	(AB017186) cardiolipin synthase [Clostridium perfringens]	(AJ002571) YkhA [Bacillus subtilis]	IN ICDC-MINE INTERGENIC REGION [Escherichia coli]	HYPOTHETICAL TRANSCRIPTIONAL REGULATOR	INTERGENIC REGION PRECURSOR [Bacillus subtilis]	subtilis]	(VEGETATIVE PROTEIN 220) (VEG220) [Bacillus	ALPHA SUBUNIT (S COMPLEX, 42 KD SUBUNIT)	PYRUVATE DEHYDROGENASE E1 COMPONENT,	OCTAPRENYLTRANSFERASE) [Bacillus subtilis]	OCTAPRENYLTRANSFERASE (DHNA-	PROBABLE 1,4-DIHYDROXY-2-NAPHTHOATE	cereus]	(AF053927) probable spore germination protein F [Bacillus	NorA [Bacillus subtilis]	(Z82987) unknown similar to quinolon resistance protein	(Z98682) YlbM protein [Bacillus subtilis]	(Z98682) YlbN protein [Bacillus subtilis]	(299117) similar to folate metabolism [Bacillus subtilis]	[Clostridium pasteurianum]	hypothetical protein 4 - Clostridium pasteurianum	(Z99120) similar to hypothetical proteins [Bacillus subtilis]	(AF041468) ompR homolog [Guillardia theta]	(D10594) chitinase D precursor [Bacillus circulans]	(Y09322) membrane bound protein LytR [Bacillus cereus]	(AJ000325) putative membrane protein [Lactococcus lactis]	(AF065404) pXO1-97 [Bacillus anthracis]	(ENDODEOXYRIBONUCLEASE IV) [Bacillus subtilis]	NCBI gi description	

Table 1

							•	•									Z = Z
186 187	185	184	183	183	182	181	180	179	178	177	176	175	174	173	172	171	NO SEQ
Bt1Gc335 Bt1Gc336	Bt1Gc334	्रे Bt1Gc333	Bt1Gc332	Bt1Gc332	Bt1'Gc329	Bt1Gc328	Bt1Gc330	Bt1Gc327	Bt1Gc324	Bt1Gc321	Bt1Gc322	Bt1Gc317	Bt1Gc319	Bt1Gc315	Bt1Gc316	Bt1Gc309	Contig Id
Bt1G229 Bt1G230	Bt1G228	Bt1G227	Bt1G226	Bt1G225	Bt1G224	Bt1G223	Bt1G222	Bt1G221	Bt1G220	Bt1G219	Bt1G218	Bt1G217	Bt1G216	Bt1G215	Bt1G214	Bt1G213	Gene Id
1033-782 1-400	1-549	1-381	922-634	609-1	1-428	548-1	1-243	744-542	1-318	24-762	424-1	787-1	1-506	191-1	721-1	1159-1	Position
g225559 g549496	g2633168	g548832	g1881373	g2499210	g2633471	g96807	g1945096	g2649585	g4155453	g466778	g2618995	g2127819	g1706795	g1730911	g2635792	g1075972	NCBI gi
409 154	446	114	227	684	539	509	183	124	285	841	333	261	374	212	235	806	nap s
409 200	493	113	245	698	560	561	192	179	257	698	336	363	332	228	297	880	BlastP Score
3.50E-38 4.90E-16	4.40E-47	1.80E-13	8.30E-21	8.20E-69	3.50E-54	1.20E-53	1.70E-14	1.00E-13	1.30E-21	8.20E-69	1.90E-30	2.60E-33	5.00E-30	1.10E-18	7.50E-26	4.30E-88	BlastP- Prob
98 34	51	29	49	67	73	58	54	45	48	63	46	31	45	69	28	46	% Ident
17 65	58	76	59	97	34	20	19	24	23	50	85	94	43	17	47	59	%Cvrg
[Bacillus subtilis] ORF IS231C [Bacillus thuringiensis] HYPOTHETICAL 22.1 KD PROTEIN IN HEML-PFS	[Pseudomonas aeruginosa] (Z99108) similar to iron(III) dicitrate transport permease						[Archaeoglobus fulgidus] (D88802) S. lividans chloramphenicol resistance protein;	[Helicobacter pylori J99] (AE001034) ABC transporter, ATP-binding protein	(AE001517) putative SODIUM/ALANINE SYMPORTER	subtilis] (M89774) lysine specific permease [Escherichia coli]	Methanococcus jannaschii [Methanococcus jannaschii] (AF027868) putative alanine acetyl transferase [Bacillus	FHUB [Bacillus subtilis] cobalt transport ATP-binding protein O homolog -	INTERGENIC REGION [Bacillus subtilis] FERRICHROME TRANSPORT PERMEASE PROTEIN	HYPOTHETICAL 42.6 KD PROTEIN IN BSAA-ILVD	perfringens] (Z99120) similar to oligoendopeptidase [Bacillus subtilis]	amyloli tetB(P)	NCBI gi description

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(Z73234) YneT [Bacillus subtilis]	57 36	1.80E-09	138	104	g1405460	605-442	Bt1G254	Bt1Gc366	208
MAJOR COLD SHOCK PROTEIN CSPA [Bacillus cereus]	99 99	1.00E-31	348	348	g2493761	10-210	Bt1G253	Bt1Gc367	207
(L77763) neutral protease A [Bacillus thuringiensis]	93 8	2.20E-09	147	178	g2239276	576-447	Bt1G252	Bt1Gc368	206
INTERGENIC REGION [Bacillus subtilis]					(
HYPOTHETICAL 22.9 KD PROTEIN IN NUCB-AROD	27 63	4.20E-10		119	g1730977	1-397	Bt1G251	Bt1Gc368	206
SMF PROTEIN [Bacillus subtilis]	32 28	9.90E-08	127	67	g3915864	413-1	Bt1G250	Bt1Gc365	205
PROTEIN) [Pseudomonas syringae]					(-,	
ACETYLTRANSFERASE (TABTOXIN RESISTANCE	33 99	6.10E-25		237	g136472	959-432	Bt1G249	Bt1Gc364	204
[Bacillus subtilis] (AB015998) CBP21 precursor [Serratia marcescens]	53 29	3.80E-09	135	138	g3308998	179-1	Bt1G248	Bt1Gc364	204
penicilin binding protein 5 - Bacillus subtilis (fragment)	39 64	8.30E-37	396	268	g2119798	1-697	Bt1G247	Bt1Gc362	203
(D86417) YfmL [Bacillus subtilis]				192	g2443255	1-418	Bt1G246	Bt. Gc358	202
(Z99108) yhbJ [Bacillus subtilis]	-			335	g2633223	566-130	Bt1G245	Bt1Gc357	201
(AF008930) ATPase [Bacillus subtilis]				544	g2293447	1-554	Bt1G244	Bt1Gc349	200
EXODEOXYRIBONUCLEASE [Bacillus subtilis]	54 · 23	2.20E-13		140	g585113	584-273	Bt1G243	Bt1Gc356	199
THIS ENTRY (YDFS_BACSU). [Bacillus subtilis]								•	
(AB001488) SIMILAR TO YDFS GENE PRODUCT OF	24 94	1.50E-12	167	111	g1881359	645-1	Bt1G242	Bt1Gc353	198
(U01945) streptothricine-acetyl-transferase [Campylobacter	36 99	1.00E-31	348	307	g732998	711-168	Bt1G241	Bt1Gc320	197
(U81516) unknown [Myxococcus xanthus]	44 60	2.90E-25		285	g2897873	674-285	Bt1G240	Bt1Gc352	196
(U75904) ProP [Escherichia coli]	52 28	1.10E-39		366	g1661219	431-1	Bt1G239	Bt1Gc351	. 195
(L80006) indolepyruvate decarboxylase [Erwinia herbicola]	46 40	8.10E-46		484	g1507711	684-1	Bt1G238	Bt1Gc350	. 194
DEHYDROGENASE 2 [Bacillus subtilis]									
PROBABLE NADH-DEPENDENT BUTANOL	67 16	3.60E-18	224	208	g3023263	517-706	Bt1G237	Bt1Gc347	193
(Z99114) similar to hypothetical proteins [Bacillus subtilis]	64 74	3.10E-30		424	g2634307	1-394	Bt1G236	Bt1Gc347	193
DEHYDROGENASE 1 [Bacillus subtilis]					:			k. *	
PROBABLE NADH-DEPENDENT BUTANOL	67 35	1.30E-47	498	461	g3023262	1-410	Bt1G235	Bt1Gc342	192
INTERGENIC REGION (Escherichia coli)				010	8170000	010-02	7040110	5170071	5
HADOLHELICVI 38 & KD BBOLEINI IN EVBB WEDV			_	616	6140635	813-82	B+1G234	Br1G6341	101
(101566) 13 8 bd OBE [Plasmid ColE1]	38 42			13.5	6144317	60-313	B+1G233	Br1Gc340	190
(X57583) mccE (Escherichia coli)	60 41	4 00E-30	417	41.7	0682770	548-065	B+1G232	B+1G-330	180
INTERGENIC REGION [Bacillus subtilis]					(
INTERGENIC REGION [Escherichia coli] HYPOTHETICAL 61.8 KD PROTEIN IN GLNQ-ANSR	50 50	1.80E-71	723	695	g1731076	1-824	Bt1G231	Bt1Gc337	188
,	Ident Cyrg	Prob Ide	Score	43	((NO
NCBI gi description	, , ,	1		aat_ B	NCBI gi	Position	Gene Id	Contig Id	SEQ
		Table 1	.					A.	
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227	226		225	224		222		221			219	218		217	216			215	214		213		212	212	211					210	9	8 E	SEQ
Bt1Gc398	Bt1Gc396	Bt1Gc393	Bt1Gc393	Bt1Gc392	Bt1Gc390	Bt1Gc386		Bt1Gc384		Bt1Gc387	Bt1Gc388	Bt1Gc383	•	Bt1Gc382	Bt1Gc373	.··.	•	Bt1Gc379	Bt1Gc380		Bt1Gc377	÷.	Bt1Gc376	Bt1Gc376	Bt1Gc374				51100072	B+1G-272	D+10-370	Contig to	
Bt1G277	Bt1G276	Bt1G274	Bt1G273	Bt1G272	Bt1G271	Bt1G270		Bt1G269		Bt1G268	Bt1G267	Bt1G266		Bt1G265	Bt1G264			Bt1G263	Bt1G262		Bt1G261		Bt1G260	Bt1G259	Bt1G258				5110607	B+1G257	D+10066	Gene Id	
1-453	432-697	879-244	174-97	679-1	1-578	570-1		1-510		671-1	642-1	791-27		1-1117	1-721			1-589	463-1		1-646		239-567	1-126	1-633				1-200	1-205	676 00	Position	
g3183519	g48//899 g549114	g2337809	g586023	g1708643	g1934809	g2274944	1	g1706797	(g2501678	g2293198	g3688826	· •	g1169000	g2226222			g3123227	g2633757	,	g3256693	(g2507017	g1945051	g4033506		•		87000077	62880830	~7676767	NCBI gi	
660	374	492	118	456	461	575		308	,	931	445	336		1178	309			693	231		139		330	152	714				. 17	130	305	nap Score	aat_
609	301 410	516	118	500	409	469		355		857	443	277		1181	369			744	288		214		263	153	754				133	150	1	Score	BlastP
2.20E-59	9.70E-27 2.70E-38	1.60E-49	2.40E-07	7.90E-48	3.50E-38	1.50E-44		1.80E-32		1.20E-85	8.70E-42	4.50E-24		5.40E-120	6.00E-34		•	1.10E-73	2.30E-25		9.30E-17		1.00E-22	2.50E-10	9.60E-75				2.000.11	3.50E-72	5 500 70	Prob	BlastP-
84	80	_	85	43	-	58		42			42	30		57	42			69	50		26		68	67	70				4			Ident	%
35	19	100	96	35	36	48		51		74	40	61		34	65			39	31		45		29	· =	73					3 6	<u>،</u>	Cyrg	%
ELEMENT IS23 IF [Bacillus thuringiensis] 5 GID PROTEIN [Bacillus subtilis]		_	STAGE 1	(US1115	_	(AJ0003					_		COLLAGENASE) [Clostridium perfringens]			SYNTHETASE) [Bacillus subtilis]	(GLUTAMINE AMIDOTRANSFERASE) (GMP	GMP SY	_	horikoshii]	_			stearothermophilus] 1 (U63928) L1 protein [Bacillus cereus]		elegans]	EMBL: C08550 comes from this gene [Caenorhabditis	dehydrogenase/reductase C-terminus: cDNA FST			(700100)	NCBI gl description	

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Table 1

(GLUCOSAMINE-6-PHOSPHATE DEAMINASE)			!	!	Ç				
GLUCOSAMINE-6-PHOSPHATE ISOMERASE		-	293	238	g3122426	918-529	Bt1G315	Bt1Gc450	260
INTERGENIC REGION [Bacillus subtilis] (AF017113) YvoA [Bacillus subtilis]	48 70	1.10E-41	442	442	g2618854	515-1	Bt1G314	Bt1Gc450	260
HYPOTHETICAL 43.3 KD PROTEIN IN QOXD-VPR	35 79	3.00E-39	419	406	g732351	1-919	Bt1G313	Bt1Gc453	259
(Z95120) hypothetical protein Rv3225c [Mycobacterium	31 ' 57	1.10E-39	423	357	g3261741	1-816	Bt1G312	Bt1Gc444	258
INTERGENIC REGION [Bacillus subtilis]									
HYPOTHETICAL 29.2 KD PROTEIN IN METS-KSGA	76 89	1.00E-93	933	933	g586875	681-1	Bt1G311	Bt1Gc441	257
(AB000617) YcdH [Bacillus subtilis]	38 94	3.30E-42	447	512	g2415736	145-1051	Bt1G310	Bt1Gc442	256
3'REGION [Bacillus subtilis]									
HYPOTHETICAL 48.3 KD PROTEIN IN KATB	75 44	2.30E-66	675	745	g1177018	314-920	Bt1G309	Bt1Gc437	255
tuberculosis]					1				
(AL022004) hypothetical protein Rv0846c [Mycobacterium	39 . 9	5.20E-06	115	94	g2916905	1-136	Bt1G308	Bt1Gc436	254
INTERGENIC REGION [Bacillus subtilis]					(
HYPOTHETICAL 53.3 KD PROTEIN IN SFP-GERKA	34 47	6.60E-27	306	263	g2829802	1-733	Bt1G307	Bt1Gc439	253
subtilis			. 6	į					ļ
TRANSCRIPTIONAL REGULATOR CTSR [Bacillus			468	489	£586897	489-28	Bt1G306	Bt1Gc438	252
bacteriolytic enzyme lytP - Bacillus sp [Bacillus sp.]		•	402	337	£2126822	485-1	Bt1G305	Bt1Gc435	251
(Z99112) penicillin-binding protein [Bacillus subtilis]	36 46	2.70E-31	344	307	g2634067	1-545	Bt1G304	Bt1Gc434	250
[Bacillus subtilis]								•	
(Z99121) similar to heavy metal-transporting ATPase	76 26	5.30E-74	747	809	g2635863	625-1	Bt1G303	Bt1Gc431	249
[Staphylococcus carnosus]									
glucose-specific, factor IIA - Staphylococcus carnosus									
phosphotransferase system enzyme II (EC 2.7.1.69),	61 14	2.90E-19	240	290	g628925	188-472	Bt1G302	Bt1Gc430	248
[Bacillus subtilis]				-					
(Z99108) similar to hypothetical proteins from B. subtilis	31 76	2.60E-08	127	96	g2633180	523-1	Bt1G301	Bt1Gc428	247
INTERGENIC REGION (F402) [Escherichia coli]									
HYPOTHETICAL 43.0 KD PROTEIN IN PROK-TAG	40 · 28	1.10E-07	129	221	g586703	338-1	Bt1G300	Bt1Gc429	246
(AF086638) CumA precursor [Pseudomonas putida GB-1]	37 35	2.50E-26	299	202	g4580028	564-1	Bt1G299	Bt1Gc425	245
maritima]									
(U67196) DNA-binding response regulator [Thermotoga	38 62	6.30E-23	265	238	g1575577	937-507	Bt1G298	Bt1Gc423	244
INTERGENIC REGION [Bacillus subtilis]							,		
HYPOTHETICAL 20.7 KD PROTEIN IN BLTR-SPOIIIC	43 99	6.00E-34	369	325	g1731125	329-965	Bt1G297	Bt1Gc422	243
,	at Cyrg	Prob Ident	Score	TD.				(NO
NCBI gi description	. · . %	•	•	aat_ l	NCBI gi	Position	Gene Id	Contig Id	SEQ
		Table 1							

Table 1

[Bacillus subtilis] 1 GMP SYNTHASE (GLUTAMINE-HYDROLYZING) (GLUTAMINE AMIDOTRANSFERASE) (GMP	84 3 41	3.00E-94	938	913	g3123227	1-628	Bt1G336	Bt1Gc472	278
	82 52	2.80E-68	693	688	g3915596	1-464	Bt1G335	Bt1Gc477	277
	40 66	9.10E-15	188	245	g2506860	599-162	Bt1G334	Bt1Gc467	276
	30 59	2.00E-11	160	105	g4115379	918-462	Bt1G333	Bt1Gc476	275
	41 41	6.10E-09	133	116	g2108269	175-1	Bt1G332	Bt1Gc476	275
7 (AJ010111) pyruvate carboxylase [Bacillus cereus]	94 17	5.40E-81	813	811	g4584148	493-1	Bt1G331	Bt1Gc475	274
1 (AF027868) YocH [Bacillus subtilis]	46 61	2.80E-29	325	316	g2619050	742-79	Bt1G330	Bt1Gc471	273
	71 75	3.10E-101	1004	1058	g2828499	845-1	Bt1G329	Bt1Gc470	272
16 (Z75208) hypothetical protein [Bacillus subtilis]		1.30E-16	209	203	g1770012	648-825	Bt1G328	Bt1Gc465	271
_	55 98	4.90E-39	417	391	g2340013	562-109	Bt1G327	Bt1Gc469	270
		;	!	!				-	;
	63 59	6.90E-88	878	860	g1731006	786-1	Bt1G326	Bt1Gc455	269
_ :	23 100	1.90E-21	178	154	g1934789	917-173	Bt1G324	Bt1Gc460	268
[Bacillus subtilis]								•	
ALPHA) (VEGETATIVE PROTEIN 63) (VEG63)					(
SUCCI	83 49	1.20E-67	687	806	g3183562	1-565	Bt1G323	Bt1Gc459	267
DNAA [Bacillus subtilis]									
	79 66	1.10E-119	1178	1219	g118704	1-890	Bt1G322	Bt1Gc458	266
			,	:	0				1
3'REGION (ORF3) [Bacillus methanolicus]	30 .	8 40F-17	160	144	0141355	660-362	Bt1G321	Rt1Gc446	265
	53 93	1.20E-28	319	308	g732151	1-335	Bt1G320	Bt1Gc446	265
DEAMINASE) [Bacillus subtilis]									
[Bacillus cereus] 6 PROBABLE D-SERINE DEHYDRATASE (D-SERINE	64 36	8.50E-51	528	491	g1711354	751-262	Bt1G319	Bt1Gc452	264
_	39 37	9.00E-24	273	197	g4584112	145-510	Bt1G318	Bt1Gc449	263
		1.70E-27	308	,251	g1945654	152-643	Bt1G317	Bt1Gc451	262
					•				
6 (AE000933) dTDP-glucose 4,6-dehydratase	50 36	1.10E-32	357	317	g2622920	600-238	Bt1G316	Bt1Gc443	261
NCBI gi description	=		Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	8 8
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122	g1124822 1064 g2811132 1046	1064 1074 1046 1011	1064 1046	1064 1074 1046 1011
g4753857	121		156	156 6.70E-11 27
g538771	779	779 795		795
g112703	619	619 533		533
g118721	406		258	258 2.70E-21 74
g2833192	307	307 328		328
g1730917	507		536	536 1.20E-51 53
g2266425	638		645	645 3.40E-63 72
g2635858	339	339 442		442
g416901	981	981 973		973
g141422	106	106 119		119
g548931	1161	1161 579		579
;				
	77	77 139		139
g1724012 145	S	5 227		227 1
g1176771 266	9	6 212		212
g1176770 5:	554	54 549		549
	448	48 454	-	454
	801	_	694	694 2.20E-68 75
g4584128 1	120	20 136	_	136
NCBI gi n	nap Score	at_ BlastP nap Score core	BlastP BlastP- Score Prob	BlastP Score

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coli] 50 INSERTION SEQUENCE IS232 PUTATIVE ATP-	97 5	1.20E-62	640	638	g2497392	901-523	Bt1G403	Bt1Gc553	329
	43 21	3.40E-33	372	339	g728913	615-1	Bt1G402	Bt1Gc550	328
13 HYPOTHETICAL 34.0 KD PROTEIN IN COTF-TETB	· :	9.50E-14	181	165	g586817	661-784	Bt1G401	Bt1Gc551	327
	··	5.20E-44	46	451	g3308998	1-440	Bt1G400	Bt1Gc551	327
13 (D86346) crystal protein [Bacillus thuringiensis]	43 1	6.90E-19	238	184	g1434920	1-310	Bt1G399	Bt1Gc549	326
76 (AF027868) YoaR [Bacillus subtilis]	31 7	5.30E-26	294	250	g2619029	876-185	Bt1G398	Bt1Gc548	325
nucleopolyhedrovirus] 60 (AF008220) YtaB [Bacillus subtilis]	56 . 6	4.00E-21	248	272	g2293285	1-279	Bt1G397	Bt1Gc547	324
[Bacillus subtilis] 36 (AF081810) viral enhancing factor 1 [Lymantria dispar	23 3	8.60E-23	274	115	g3822300	1-847	Bt1G396	Bt1Gc546	323
	42 31	3.30E-41	443	338	g1881236	1-711	Bt1G395	Bt1Gc544	322
47 (Y14081) hypothetical protein [Bacillus subtilis]	51 4	3.00E-16	202	182	g2226190	845-640	Bt1G394	Bt1Gc543	321
[Synechocystis sp.] 92 (U66480) YncE [Bacillus subtilis]	40 9	8.40E-28	311	282	g1750129	423-1	Bt1G393	Bt1Gc543	321
cereus] 62 HYPOTHETICAL 33.3 KD PROTEIN SLL1263	33 6	1.60E-24	280	282	g2501575	451-1019	Bt1G392	Bt1Gc539	320
[Enterobacter cloacae] 6 (AF053927) probable spore germination protein F [Bacillus	79 : 46	8.70E-10	141	125	g2984723	1-100	Bt1G391	Bt1Gc539	320
(INDOLEPYRUVATE DECARBOXYLASE)	· ·.								
INDOL	50 41	8.00E-55	566	554	g118333	682-1	Bt1G390	Bt1Gc542	319
99 (Z99121) similar to hypothetical proteins [Bacillus subtilis]	63	1.40E-32	356	351	g2635865	489-187	Bt1G389	Bt1Gc522	318
14 HYPOTHETICAL 39.9 KD PROTEIN IN AMYLASE	32 1	6.00E-06	112	. 95	g418336	676-532	Bt1G388	Bt1Gc540	317
38 XANTHINE PERMEASE [Bacillus subtilis]		3.10E-46	485	533	g1172036	1-501	Bt1G387	Bt1Gc540	317
(ENTE	٠.	1.100		Š	9000				
DNA ALKYLTRANSFERASE) [Bacillus subtilis] 78 HEMOI VSIN BI BINIDING COMPONENT DRECTIDECOR	%	1 10E-103	1027	990	a2507017	881 <u>-</u> 1	Br1G386	Br1Gc538	ગ
	44 93	2.60E-33	363	337	g113335	381-880	Bt1G385	Bt1Gc537	315
ALKYLTRANSFERASE [Bacillus subtilis]	. •							 - -	!
[Cucumis sativus] 1 METHYLPHOSPHOTRIESTER-DNA	52 61	2.00E-35	383	377	g113333	1-385	Bt1G384	Bt1Gc537	315
4 (M16219) glyoxysomal malate synthase (EC 4.1.3.1)	39 74	0.0041	78	118	g167521	769-975	Bt1G383	Bt1Gc533	314
NCBI gi description	% % Ident Cvrg	BlastP- Prob I	BlastP Score	aat_ nap Score	NCBI gi	Position	Gene Id	Contig Id	NO SEQ

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beta-lac [Bacill	92 11	1.40E-10	153	137	g80073	1443-1338	Bt1G423	Bt1Gc572	345
19 (AB001488) SIMILAR TO YQFU, YXKD, YITB OF B. SURTILIS [Bacillus subtilis]	44 1	5.50E-06	111	97	g1881335	161-1	Bt1G422	Bt1Gc572	345
	35 99	7.80E-09	132	114	g2633659	266-44	Bt1G421	Bt1Gc569	344
	37 18	7.50E-09	141	65.	g266724	621-374	Bt1G420	Bt1Gc541	343
INTERGENIC REGION (ORF3) [Bacillus subtilis] 9 PROBABLE RIBOSOMAL PROTEIN IN NUSA-INFB INTERGENIC REGION (ORF4) [Bacillus subtilis]	61 99	1.50E-28	318	309	g418462	418-717	Bt1G419	Bt1Gc570	342
	59 99	1.50E-21	252	290	g418461	148-420	Bt1G418	Bt1Gc570	342
		1.90E-06	117	85	g2634032	1-136	Bt1G417	Bt1Gc570	342
3 codV protein - Bacillus subtilis []	52 63	9.30E-45	471	534	g2126912	458-1026	Bt1G416	Bt1Gc568	341
0 GID PROTEIN [Bacillus subtilis]	58 30	1.40E-34	375	340	g3183519	1-395	Bt1G415	Bt1Gc568	341
	80 58	8.50E-138	1349	1418	g3183185	1-1074	Bt1G414	Bt1Gc562	340
Jannaso (AL03	28 17	2.40E-07	128	74	g3451437	600-869	Bt1G413	Bt1Gc567	339
	35 61	2.30E-18	222	178	g3024926	665-1096	Bt1G412	Bt1Gc565	338
					(
		1.30E-13	177	173	g141373	775-515	Bt1G411		337
[Bacillus subtilis] 0 (D29673) hydrolase [Lactobacillus sp.]	27 .30	1.20E-11	168	<u>چ</u>	g473953	503-1	Bt1G410	Bt1Gc559	336
_	51 70	2.20E-75	760	746	g1945088	920-115	Bt1G409	Bt1Gc560	335
	•	•			. (
thermocatenulatus] 2 HYPOTHETICAL 42.6 KD PROTEIN IN BSAA-ILVD	62 . 8 2	3.70E-89	890	924	g1730911	1148-233	Bt1G408	Bt1Gc557	334
	44 78	6.50E-69	699	564	g1321706	1072-91	Bt1G407	Bt1Gc555	333
iugidusj 3 (AB014075) Orf5u [Clostridium histolyticum]	30 63	2.90E-34	372	320	g3868868	1-889	Bt1G406	Bt1Gc556	332
	24 76	2.00E-19	232	140	g2650107	833-165	Bt1G405	Bt1Gc554	331
BINDING PROTEIN [Insertion sequence IS232] 5 ORF IS231C [Bacillus thuringiensis]	99 15	7.70E-34	368	368	g225559	391-612	Bt1G404	Bt1Gc552	330
• •	% % Ident Cvrg	BlastP- Prob L	BlastP Score	aat_ nap Score	NCBI gi	Position	Gene Id	Contig Id	SEQ NO

Table 1

SEQ C			348 B	. 349 В			351 B	352 B	353 B	354 B	355 B	356 B	357 B	357 B	358 B	358 B	359 B	360 1
Contig Id	Bt1Gc574	Bt1Gc573	Bt1Gc571	Bt1Gc577	10,575	מנוטנטיט	Bt1Gc578	Bt1Gc566	Bt1Gc579	Bt1Gc583	Bt1Gc584	Bt1Gc585	Bt1Gc582	Bt1Gc582	Bt1Gc590	Bt1Gc590	Bt1Gc588	Bt1Gc591
Gene Id	Bt1G424	Bt1G425	Bt1G426	Bt1G427	B+1C428	0740110	Bt1G429	Bt1G430	Bt1G431	Bt1G432	Bt1G433	Bt1G434	Bt1G435	Bt1G436	Bt1G437	Bt1G438	Bt1G439	Bt1G440
Position	1386-101	910-1	1-762	1-1424	1016-1	1-0101	361-1	538-38	1-1009	839-1	252-1031	1072-132	1-165	278-1101	1-275	289-540	1-798	480-1
NCBI gi	g2094843	g1354814	g1346893	g1731087	G301 < 000	83713770	g3123299	g3915543	g3322837	g2507445	g2661703	g732405	g1575578	g2634226	g2634266	g2634267	g121467	g2909586
nap Score	711	229	226	185	305	373	572	566	196	881	138	105	114	862	267	308	1081	98
BlastP Score	499	356	154	233	376	0/0	480	566	375	877	231	190	148	875	276	322	420	137
BlastP- Prob	1.00E-47	1.40E-32	2.40E-14	1.60E-19	1 405-34	1.40E-34	1.00E-45	8.00E-55	4.70E-34	8.90E-88	2.50E-19	4.50E-14	1.00E-09	1.40E-87	4.30E-24	5.70E-29	1.60E-71	2.30E-09
% Ident	39	29	28	37	3,	J.	8	62	27	61	27	31	53	59	62	76	81	26
%Cvrg	89	79	28	4	, ,	ę	26	99	58	63	82	24	13	83	57	. 99	63	76
NCBI gi description	(Z95324) mgtE [Mycobacterium tuberculosis]	(U57065) ZmaR [Bacillus cereus]	PTS SYSTEM, GLUCOSE-SPECIFIC IIABC COMPONENT (EIIABC-GLC) (GLUCOSE-PERMEASE IIABC COMPONENT) (PHOSPHOTRANSFERASE	ENZYME II, ABC COMPONENT) (EII-GLC / EIII-GLC) [Mycoplasma genitalium] HYPOTHETICAL 39.0 KD PROTEIN IN GLNQ-ANSR	INTERGENI© REGION [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL TRANSPORT PROTEIN IN NDHF- CSGA INTERGENIC REGION (ORF1) [Bacillus subtilis]	HYPOTHETICAL 18.9 KD PROTEIN IN CYPA-AADK	(AE001229) T. pallidum predicted coding region TP0544	D-SERINE DEHYDRATASE (D-SERINE DEAMINASE) [Escherichia coli]	(AL009204) hypothetical protein SC9B10.17 [Streptomyces coelicolor]	HYPOTHETICAL TRANSCRIPTIONAL REGULATOR IN FEUA-SIGW INTERGENIC REGION (ORF3) [Bacillus substitic]	(U67196) histidine protein kinase [Thermotoga maritima]	(Z99113) similar to alcohol dehydrogenase [Bacillus subtilis]	(Z99114) yoaS [Bacillus subtilis]	(Z99114) similar to transcriptional regulator [Bacillus subtilis]	PROTON/SODIUM-GLUTAMATE SYMPORT PROTEIN (GLUTAMATE-ASPARTATE CARRIER PROTEIN)	(AL021927) hypothetical protein Rv0158 [Mycobacterium

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378 Bt1	377 Bt1	376 Bt1	375 Bt1	374 Bt1		374 Bt1	373 Bt1			372 Bt1	371 Bt1	370 Bt1		369 Bt1		368 Bt1	367 Bt1	366 Bt1	365 Bt1		364 Bt1	363 Bt1	363 Bt1		362 Bt1	361 Bt1	SEQ DO CO
Bt1Gc616	Bt1Gc617	Bt1Gc614	Bt1Gc612	Bt1Gc613		Bt1Gc613	Bt1Gc605			Bt1Gc611	Bt1Gc608	Bt1Gc607		Bt1Gc603		Bt1Gc606	Bt1Gc600	Bt1Gc601	Bt1Gc599		Bt1Gc597	Bt1Gc596	Bt1Gc596		Bt1Gc592	Bt1Gc593	Contig Id
Bt1G462	Bt1G461	Bt1G460	Bt1G459	Bt1G458		Bt1G457	Bt1G456		Bt1G455	Bt1G454	Bt1G453	Bt1G452		Bt1G451		Bt1G450	Bt1G448	Bt1G447	Bt1G446		Bt1G445	Bt1G444	Bt1G443		Bt1G442	Bt1G441	Gene Id
1083-1	486-887	1-1251	1-443	1037-588		416-1	1-567		1996-1665	1545-479	853-174	247-1	•	1058-1		784-80	807-1	1-618	305-724		839-20	534-715	1-431		1182-336	1030-35	Position
g1171759	g4154073	g1175719	g586898	g400755		g400802	g4155188	C	g1731305	g2116973	g2619012	g3322943		g628966		g1168824	g3914084	g548909	g2633752	•	g586703	g2635584	g2577968		g2621332	g3861147	NCBI gi
249	240	758	312	248		521	586		174	1253	79	215		509		864	518	519	177		585	80	281		187	199	nap S
452	265	799	292	260		520	532		131	1289	119	247		544		588	545	532	248		510	108	241		234	215	BlastP Score
9.60E-43	6.30E-23	1.60E-79	8.70E-26	2.10E-22		6.00E-50	3.20E-51		1.00E-08	1.90E-131	1.10E-06	2.90E-20		1.70E-52		3.70E-57	1.90E-52	3.20E-51	4.00E-21		6.90E-49	2.70E-06	2.20E-20		1.20E-19	4.30E-17	BlastP- Prob
31	42	40	41	36		68	58		45	67	49	51		40		65	40	52	38		41	38	37		34	27	% Ident (
82	99	2	80	89		55	4	1	88	<u>1</u> 00	14	. 16		49		100	34	48	99		8	35	%	•	99	60	Cvrg
BACILLOLYSIN PRECURSOR (NEUTRAL PROTEASE) [Bacillus brevis]	subtilis] (AL035161) conserved hypothetical protein [Streptomyces	PENICILLIN-BINDING PROTEIN 3 (PBP 3) [Bacillus	HYPOTHETICAL 21.0 KD PROTEIN IN LYSS-MECB	PETP PROTEIN [Rhodobacter capsulatus]	(PHOSPHOGLYCEROMUTASE I) (PGAM I) (BPG-	[Helicobacter pylori J99] PHOSPHOGLYCERATE MUTASE 1	(AE001495) short-chain fatty acids transporter	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 14.7 KD PROTEIN IN IDH-DEOR	(D87979) YfnK [Bacillus subtilis]	(AF027868) fatty acid desaturase [Bacillus subtilis]	(AE001239) lysyl-tRNA synthetase (lysS-1) [Treponema	plasmid pDB101 [Streptococcus pyogenes]	hypothetical protein gamma - Streptococcus pyogenes	[Bacillus subtilis]	CYTOCHROME C-TYPE BIOGENESIS PROTEIN CCDA	MUTS2 PROTEIN [Bacillus subtilis]	SERINE TRANSPORTER [Escherichia coli]	(Z99111) ykvS [Bacillus subtilis]	INTERGENIC REGION (F402) [Escherichia coli]	HYPOTHETICAL 43.0 KD PROTEIN IN PROK-TAG	(Z99119) yual [Bacillus subtilis]	(Y15254) SapB protein [Bacillus subtilis]	thermoautotrophicum]	(bcr1) [kickettsia prowazekii] (AE000813) conserved protein [Methanobacterium	(AJ235272) BICYCLOMYCIN RESISTANCE PROTEIN	NCBI gi description

Table 1

acetyltransferase; P44305 (189) [Bacillus subtilis] (AF015775) carboxypeptidase [Bacillus subtilis]	40 34	3.70E-07	121	120	ę2415396	1-1141	Bt1G485	Bt1Gc640	396
	26 99	2.00E-10	147	78	g1945109	288-756	Bt1G484	Bt1Gc640	396
(Y14084) hypothetical protein [Bacillus subtilis]	42 .78	1.90E-60	619	616	g2226258	1-1010	Bt1G483	Bt1Gc635	395
	91 9	1.10E-14	195	195	g225559	867-739	Bt1G482	Bt1Gc637	394
(Z82015) yukJ [Bacillus subtilis]	41 37	7.00E-08	123	85	g1665854	223-382	Bt1G481	Bt1Gc632	393
	38 100	2.00E-42	449	395	g2635861	195-845	Bt1G480	Bt1Gc636	. 392
ORF IS231C [Bacillus thuringiensis]	50 2,1	7.00E-17	215	129	g225559	1049-686	Bt1G479	Bt1Gc634	391
(Z94043) hypothetical protein [Bacillus subtilis]	35 100	8.90E-40	424	375	g1945654	520-1285	Bt1G478	Bt1Gc610	390
cereus]	٠.								
(AJ011526) D-stereospecific peptide hydrolase [Bacillus	70 88	1.50E-122	1205	1277	g4127525	1-1038	Bt1G477	Bt1Gc630	389
INTERGENIC REGION [Bacillus subtilis]									
HYPOTHETICAL 21.7 KD PROTEIN IN NRGB-SPOIIQ	35 74	5.80E-20	237	256	g3025311	1-433	Bt1G476	Bt1Gc631	388
ELEMENT IS231E [Bacillus thuringiensis]									
TRANSPOSASE FOR INSERTION SEQUENCE	72 21	7.50E-26	296	337	g549113	1199-904	Bt1G475	Bt1Gc629	387
tuberculosis]	: .								
_	46 : 22	3.90E-07	121	132	g2896709	1-182	Bt1G474	Bt1Gc629	387
INTERGENIC REGION [Bacillus subtilis]	٠.								
HYPOTHETICAL 43.8 KD PROTEIN IN XPAC-ABRB	25 .37	1.80E-08	136	134	g586865	1-434	Bt1G473	Bt1Gc628	386
[Bacillus firmus]	. ,								
(AF084104) maltose transportor ATP-binding protein	53 33	1.00E-31	348	325	g3688811	926-559	Bt1G472	Bt1Gc620	385
(AF119621) Ditl [Pseudomonas abietaniphila]	29 · 63	2.00E-12	168	176	g4455080	1-926	Bt1G471	Bt1Gc620	385
PUTATIVE ARSENATE REDUCTASE [Bacillus subtilis]	77 73	3.00E-39	419	409	g1168520	252-563	Bt1G470	Bt1Gc627	384
INTERGENIC REGION [Bacillus subtilis]	٠.				•				
HYPOTHETICAL 38.3 KD PROTEIN IN CWLA-CISA	85 21	4.40E-22	257	306	g1176771	1-223	Bt1G469	Bt1Gc627	384
	36 ∴ 57	3.30E-17	211	171	g2497400	//8-402	Bt1G468	R110c625	383
	٠.		•	į			}		
HYPOTHETICAL 43.7 KD PROTEIN IN KATB	61 · 10	6.90E-09	140	118	g1177016	529-651	Bt1G467	Bt1Gc624	382
(D90917) hypothetical protein [Synechocystis sp.]	33 82	2.00E-19	232	.227	g1653921	899-422	Bt1G466	Bt1Gc622	381
3'REGION [Bacillus subtilis]								-	
	39 94	5.20E-51	530	441	g1723610	74-926	Bt1G465	Bt1Gc623	380
HI1721	. •				,				
	38 60	4.10E-19	229	184	g2497400	919-484	Bt1G464	Bt1Gc619	379
(L77763) neutral protease A [Bacillus thuringiensis]	32 63	4.00E-37	399	264	g2239276	1083-1	Bt1G463	Bt1Gc616	378
Men gi aescription	Ident Cvrg	Prob Id	Score	nap Score	19 TO 19	1 0310101	Celle	6 10	8
NCRI of description	%	BlastP-	BlastP		NCRI oi	Position	Gene Id	Contig Id	SEQ
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28 LIPASE 1 PRECURSOR (TRIACYLGLYCEROL LIPASE) [Psychrobacter immobilis] 51 PUTATIVE NAD(P)H NITROREDUCTASE YFKO [Bacillus subtilis]	8 31 3 58	5.40E-08 2.00E-33	130 364	65 3 <i>5</i> 7	g1346454 g3915460	1-417 335-1	Bt1G504 Bt1G505	Bt1Gc652 Bt1Gc653	411 412
13	_	1.10E-10		126	g2507017	1336-1194	Bt1G503		410
32		1.80E-07		57	g2127290	915-676	Bt1G502		410
ა:	0	1.90E-110	1091	1086	g231698	721-1	Bt1G501	Bt1Gc657	409
2	2 59	7.90E-32	349	313	g586039	678-346	Bt1G500	Bt1Gc656	408
72	53	2.60E-49	514	509	g1345692	1-473	Bt1G499	Bt1Gc654	407
76	9 43	3.80E-09	135	113	g3256746	262-1	Bt1G498	Bt1Gc655	406
28	6 49	6.20E-16	199	173	g2688589	1138-919	Bt1G497	Bt1Gc647	405
100	49 :	3.30E-65	664	679	g2624002	948-109	Bt1G496	Bt1Gc649	404
78		1.10E-09	140	200	g1723608	21-1317	Bt1G495	Bt1Gc650	403
65	1 57	1.00E-31		402	g3599371	442-1	Bt1G494	Bt1Gc643	· 402
&	_	4.60E-20	238	228	g2635234	766-391	Bt1G493	Bt1Gc648	. 401
99	5 62	1.30E-15	196	265	g2635235	94-357	Bt1G492	Bt1Gc648	401
					• •				
6	6 86	9.10E-06	111	111	g3122932	Jan-66	Bt1G491	Bt1Gc648	401
99	3 44	· 1.10E-43	461	435	g2226164	616-1	Bt1G490	Bt1Gc646	400
··	ب ب 1	4.00E-04	. 6	120	81170202	70/2	7070		,
N		4 90E 6	637	3	~1176707	1 700	D+1C/190	D+1C2645	200
46	7 98	9.90E-57	584	570	g2497392	968-625	Bt1G488	Bt1Gc644	398
37	6 29	9.10E-16	202	141	g2127182	1-396	Bt1G487	Bt1Gc644	398
48	9 59	5.90E-59	605	604	g1705428	953-286	Bt1G486	Bt1Gc642	397
`} %	% % Ident Cvrg	BlastP- Prob	BlastP Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	S E S
)

Table 1

426	426	425	424		424		423	422		422		421		420	419	418		418		417	416		416		415	414	414	413		413	412	NO	E SEC	250
Bt1Gc681	Bt1Gc681	Bt1Gc679	Bt1Gc676		Bt1Gc676		Bt1Gc675	Bt1Gc674		Bt1Gc674		Bt1Gc673	•	Bt1Gc672	Bt1Gc671	Bt1Gc670		Bt1Gc670		Bt1Gc666	Bt1Gc665		Bt1Gc665		Bt1Gc664	Bt1Gc662	Bt1Gc662	Bt1Gc661		Bt1Gc661	Bt1Gc653		Contig Id	
Bt1G527	Bt1G526	Bt1G525	Bt1G524		Bt1G523		Bt1G522	Bt1G521		Bt1G520		Bt1G519		Bt1G518	Bt1G517	Bt1G516		Bt1G515		Bt1G514	Bt1G513		Bt1G512		Bt1G511	Bt1G510	Bt1G509	Bt1G508		Bt1G507	Bt1G506		Gene Id	
463-748	1-341	965-621	1-752		1-756		1-1038	448-1218		26-280		1060-1		727-1014	1-548	1586-469		393-16		11-927	759-455		1-759		1-1090	1134-765	1-201	948-684	•	462-1	485-838		Position	
g2635928	g1945711	g3116222	g2149903	1	g585648	ı	g2633162	g417830		g418608		g1174634		g465650	g2340009	g1001578		g1763703		g2493595	g225559		g2497382	(g2415704	g1263187	g2226145	g116956		g1730956	g586816		NCBI gi	
274	277	140	241		500		329	814		212		93		192	463	752		193		512	490		537		<u>∞</u>	102	149	137		540	373	Score	deu Tark	
308	291	124	348		546		245	814		223		168		176	481	784		229		558	501		570		123	161	171	189		540	398	1	BlastP Score	
1.70E-27	1.10E-25	3.90E-07	8.20E-31		1.10E-52		7.60E-26	4.20E-81		1.80E-18		2.10E-12		1.70E-13	8.10E-46	6.30E-78		4.10E-19		5.60E-54	6.20E-48		4.40E-57		5.20E-07	1.40E-11	4.40E-12	7.10E-15		4.60E-52	5.10E-37	1	BlastP-	
59	49	28	33		39		25	60		48		26		38	48	39		37		38	98		85		32	30	50	47		2	61		% Ident (
23	27	. 29	33	٠.	· 35		83	86		99		60		99	4	95		99	•	92	21		29		16	43	13	. 99	*	85	92	ď	Cyre Cyre	
(Z99121) similar to maltodextrin transport system permease	(Z94043) hypothetical protein [Bacillus subtilis]	(AB007122) transporter [Arthrobacter sp.]	(U94707) penicillin-binding protein [Enterococcus faecalis]	subtilis]	PENICILLIN-BINDING PROTEIN 2B (PBP-2B) [Bacillus	subtilis]	subtilis] (Z99108) similar to multidrug resistance protein [Bacillus	DIPICOLINATE SYNTHASE, A CHAIN [Bacillus	5'REGION (ORFZ) [Bacillus subtilis]	HYPOTHETICAL 9.8 KD PROTEIN IN SPOVFA	[Haemophilus influenzae Rd]	TELLURITE RESISTANCE PROTEIN TEHB HOMOLOG	3'REGION (ORFA1) [Clostridium acetobutylicum]	HYPOTHETICAL 11.0 KD PROTEIN IN HSP18	(Z98682) YlbM protein [Bacillus subtilis]	(D64000) aspartate aminotransferase [Synechocystis sp.]	dehydratase [Bacillus subtilis]	(Z83337) similar to hydroxymyristoyl-(acyl carrier protein)	[Methanococcus jannaschii]	PUTATIVE POTASSIUM CHANNEL PROTEIN MJ0138.1	ORF IS231C [Bacillus thuringiensis]	ELEMENT IS232 [Insertion sequence IS232]	TRANSPOSASE FOR INSERTION SEQUENCE	freudenreichii subsp. shermaniil	(AJ001361) proline iminopeptidase [Propionibacterium	(U24215) HOMODA hydrolase [Pseudomonas putida]	(Y14079) hypothetical protein [Bacillus subtilis]	SPORE COAT PROTEIN D [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 21.1 KD PROTEIN IN COTD-KDUD	HYPOTHETICAL 14.7 KD PROTEIN IN COTF-TETB		NCBI gi description	

Table 1

440 Bt1Gc693 Bt1G547 779-1200 g1731066 406 417 4.9	440 Bt1Gc693 Bt1G546 1-759 g1731065 605 574 1.	439 Bt1Gc692 Bt1G545 1320-1259 g130298 86 102 6.7	439 Bt1Gc692 Bt1G544 1061-1 g1769558 787 842 4.:	437 Bt1Gc690 Bt1G542 1-256 g3023490 245 259 2.7 438 Bt1Gc689 Bt1G543 576-1 g3183496 288 267 3.9	Bt1Gc691 Bt1G541 668-1043 g3183454 381 401	435 Bt1Gc688 Bt1G539 1-606 g2632226 452 506 1.8	434 Bt1Gc687 Bt1G538 1199-628 g2648503 175 272 1.1	432 Bt1Gc677 Bt1G536 281-780 g3021327 641 480 1.0 433 Bt1Gc686 Bt1G537 281-1 g3256836 76 136 1.1	432 Bt1Gc677 Bt1G535 15-278 g131533 316 287 2.9	431 Bt1Gc685 Bt1G534 1554-619 g1730929 187 248 4.8	430 Bt1Gc684 Bt1G532 1-1072 g2293165 1264 1269 2.60 431 Bt1Gc685 Bt1G533 588-1 g3907604 271 299 1.6	Bt1Gc683 Bt1G531 1042-606 g4033455 277 254	Bt1Gc680 Bt1G529 564-1 g2612894 135 212 Bt1Gc683 Bt1G530 301-1 e1945088 261 273	NO Score Pt 427 Bt1Gc682 Bt1G528 1365-620 01620023 629 572 15
g1731066 406 417	605	g130298 86 102	g1769558 787 842	245 259 288 267	g3183454 381 401	452 506 3-19 349	g2648503 175	g3021327 641 g3256836 76	g131533 316 287	g1730929 187 248	g2293165 1264 1269 g3907604 271 299	g4033455 277 254	g2612894 135 212 g1945088 261 273	Score Score Score Score
4.90E-39 59 79 H	PI 1.10E-55 49 78 H	m 6.70E-05 95 6 1-	4.50E-84 44 75 (L	2.70E-22	60 .43	1.80E-48 46 69 (<i>A</i> 7 90E-32 37 54 (7	ho 1.10E-23 38 99 (<i>A</i> [<i>A</i>	1.00E-45 77 29 (A 1.10E-08 36 30 (A	1A 2.90E-25 72 99 PI C	C ₁ 4.80E-21 29 83 H	2.60E-129 66 56 (A 1.60E-26 36 77 (A	41 81	25 47	Prob Ident Cvrg
HYPOTHETICAL 19.7 KD PROTEIN IN GLNQ-ANSR INTERGENIC REGION [Bacillus subtilis]	PHOSPHOLIPASE C) (PI-PLC) [Bacillus thuringiensis] HYPOTHETICAL 37.1 KD PROTEIN IN BMRU-ANSR INTERGENIC REGION [Bacillus subtilis]	mortiterum] 1-PHOSPHATIDYLINOSITOL PHOSPHODIESTERASE 1-PHOSPHATIDYLINOSITOL-SPECIFIC	(U81184) phospho-beta-glucosidase [Fusobacterium	CR(VI) REDUCTASE [Pseudomonas sp.] HYPOTHETICAL 23.4 KD PROTEIN IN AAPA-SIGV INTERCENT RECTON [Bacilling subsidie]	HYPOTHETICAL 30.7 KD PROTEIN IN MCPC-KINA INTERGENIC REGION [Bacillus subtilis]	(AJ222587) YkuE protein [Bacillus subtilis]	horikoshii] (AE000962) A. fulgidus predicted coding region AF2041 [Archaeoglobus fulgidus]	(AJ005075) enzyme I [Bacillus megaterium] (AP000002) 304aa long hypothetical protein [Pyrococcus	INTERGENIC REGION [Bacillus subtilis] PHOSPHOCARRIER PROTEIN HPR (HISTIDINE- CONTAINING PROTEIN) [Bacillus subtilis]	Cps19aF [Streptococcus pneumoniae] HYPOTHETICAL 42.0 KD PROTEIN IN DAPB-PAPS	(AF008220) asparagine synthase [Bacillus subtilis] (AF094575) putative N-acetyl-mannosamine transferase	[Bacillus subtilis] SIGNAL PEPTIDASE I P (SPASE I) (LEADER	(AF015825) hypothetical glycosyl transferase [Bacillus subtilis] (D88802) S lividans olycosyl transferase: IS0636 (311)	(770580) putative ORF [Bacillus subtilis]

Table 1

461-147 g1787091 271 291 1.10E-25 54 99 1016-69 g2619050 280 313 5.20E-28 49 54 199-1 g1731007 150 152 5.90E-11 40 79 346-914 g1731008 654 508 1.10E-48 68 65 427-1 g417115 257 278 7.40E-24 41 30 839-255 g2072437 131 175 2.20E-13 30 99 538-1063 g136321 169 230 1.80E-18 30 37 204-938 g2619056 813 638 1.90E-62 63 55 1369-487 g729343 627 641 9.00E-63 44 73 1-950 g26337813 696 500 7.90E-48 66 38 1-950 g2633168 754 728 5.50E-72 48 98 1097-1290 g1706292 119 162 9.80E-12 44 22 413-1110 g3915501 770 </th
291 1.10E-25 54 313 5.20E-28 49 152 5.90E-11 40 508 1.10E-48 68 278 7.40E-24 41 175 2.20E-13 30 230 1.80E-18 30 641 9.00E-62 63 641 9.00E-63 44 347 1.30E-31 37 500 7.90E-48 66 728 5.50E-72 48 162 9.80E-12 44
1.10E-25 54 5.20E-28 49 5.20E-11 40 1.10E-48 68 7.40E-24 41 2.20E-13 30 1.80E-18 30 1.90E-62 63 9.00E-63 44 1.30E-31 37 7.90E-48 66 5.50E-72 48 9.80E-12 44

Table 1

g/033/13
5715 813
703 2.40E-69
E-09
84 (299120) similar to Na+/nucleoside cotransporter [Bacillus
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(Z99120) similar to Na+/nucleoside cotrans

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APYRIMIDINIC SITE) LYASE) [Bacillus subtilis]	\$		7 30F-77	774	741	0418587	1-810	Br1G612	Rt1Gc747	490
INTERGENIC REGION (O162) [Escherichia coli] 100 PROBABLE ENDONUCLEASE III (DNA-(APURINIC OR	77 10		3.10E-85	853	908	g729418	945-293	Bt1G611	Bt1Gc755	489
56 HYPOTHETICAL 18.8 KD PROTEIN IN GNTR-GGT	36 . :		4.50E-13	172	169	g1176283	1-275	Bt1G610	Bt1Gc754	488
97 (U58864) orf1 [Bacillus subtilis]	45		4.50E-100	993	928	g1381682	1310-1	Bt1G609	Bt1Gc753	487
57 (AB007122) transporter [Arthrobacter sp.]	34 :		5.80E-36	388	407	g3116222	422-1100	Bt1G608	Bt1Gc748	486
(PEPTIDASE D) [Escretiona con] 31 (D90899) regulatory protein PchR [Synechocystis sp.]	43		2.30E-18	222	195	g1651667	1-302	Bt1G607	Bt1Gc748	486
ALANYL-HISTIDINE DIPEPTIDASE) (CARNOSINASE)	•									
18 AMINOACYL-HISTIDINE DIPEPTIDASE (XAA-HIS		[4 4 <u>1</u>	3.80E-14	190	173	g129799	260-1	Bt1G606	Bt1Gc752	485
[Lactococcus lactis] 54 (AF036967) putative response regulator [Lactobacillus	63	_	3.60E-36	390	377	g4104605	844-1209	Bt1G605	Bt1Gc751	484
99 HYPOTHETICAL PROTEIN IN PEPC 5'REGION (ORF 2)		6 .40	1.10E-36	395	338	g732240	637-41	Bt1G604	Bt1Gc751	484
norikosniij 63 (D50098) multidrug transporter [Bacillus subtilis]	35		1.80E-48	506	536	g1856977	1-973	Bt1G603	Bt1Gc750	483
[Bacillus subtilis] 99 (AP000006) 166aa long hypothetical protein [Pyrococcus	24 5	-	6.10E-09	133	69	g3258042	102-587	Bt1G602	Bt1Gc739	482
subtilis] 23 PYRIMIDINE-NUCLEOSIDE PHOSPHORYLASE	8		1.30E-38	413	378	g730287	683-979	Bt1G601	Bt1Gc746	481
54 (Z99124) pyrimidine-nucleoside transport protein [Bacillus		080	4.80E-80	804	867	g2636487	1-643	Bt1G600	Bt1Gc746	481
epidermidis] 12 (AF009352) ATPase [Bacillus subtilis]		0 60	1.90E-10	154	138	g2271389	1137-1278	Bt1G599	Bt1Gc744	480
68 (U77778) putative membrane protein [Staphylococcus		7 31	7.60E-27	302	187	g2196513	1-1086	Bt1G598	Bt1Gc744	, 480
98 HEMOLYSIN BL BINDING COMPONENT PRECURSOR		86 98	2:20E-1/8	1/32	1829	/10/0c78	1-1101	ואנוטאו	B(10c/41	4/9
			1.10E-27	310	308	g2340007	1372-1025	Bt1G596	Bt1Gc742	478
(Z98682)	_	_	2.10E-77	779	785	g2340008	1020-1	Bt1G595	Bt1Gc742	478
(Z99110	_	-	1.40E-34	375	318	g2633511	183-843	Bt1G594	Bt1Gc740	477
DIPEPTIDYL-PEPTIDASE) (X-PROLYL-DIPEPTIDYL AMINOPEPTIDASE) (X-PDAP) [Lactococcus lactis]										
protein [Streptomyces coelicolor] 35 XAA-PRO DIPEPTIDYL-PEPTIDASE (X-PRO		42	2.70E-54	561	451	g118908	804-1	Bt1G593	Bt1Gc738	476
nCBI gi description	% cvrg	% Ident	BlastP- Prob	BlastP Score	aat_ nap Score	NCBI gi	Position	Gene Id	Contig Id	SEQ NO

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SEQ ID Contig Id NO	Gene Id	Position	NCBI gi	nap Score	BlastP Score	BlastP- Prob	% % Ident Cvrg	% Vrg	NCBI gi description
491 Bt1Gc756	Bt1G613	1-714	g3121979	459	494	3.40E-47	43	84	INTERGENIC REGION (ORFQ) [Bacillus subtilis] D-ALANINE AMINOTRANSFERASE (D-ASPARTATE
	•								AMINOTRANSFERASE) (D-AMINO ACID AMINOTRANSFERASE) (D-AMINO ACID
491 Bt1Gc756	Bt1G614	813-1242	g2499210	8	135	3.80E-09	40	32	TRANSAMINASE) [Bacillus subtilis] PUTATIVE NAD(P)H NITROREDUCTASE YDGI
				3	3		:		[Bacillus subtilis]
492 Bt1Gc/5/	B+10-615	608-1	6730175 16/78678	222	339	6.90E-33	7 <u>4</u>	4 6	(AEUUU67U) hypothetical protein [Aquitex aeolicus]
			0	9	Ş	0.100			(VEG286B) [Bacillus subtilis]
493 Bt1Gc759	Bt1G617	1229-689	g2126915	754	770	1.90E-76	83	39	codX protein - Bacillus subtilis []
494 Bt1Gc760	Bt1G618	945-131	g4514323	779	820	9.70E-82	53	74	(AB013367) unknown [Bacillus halodurans]
495 Bt1Gc761	Bt1G619	895-1	g114116	899	856	1.50E-85	57	82	ARGININE/ORNITHINE ANTIPORTER [Pseudomonas
495 Bt1Gc761	Bt1G620	1817-1127	g4155319	400	441	1.40E-41	39	41	(AE001506) putative osmoprotection binding protein
496 Bt1Gc763	Bt1G621	1018-1	g1805447	254	322	5.70E-29	25	93	(D50453) response-regulator aspartate phosphatase C
496 Bt1Gc763	Bt1G622	1018-1	g1648857	202	243	1.90E-20	22	93	(Z81356) aspartyl-phosphate phosphatases [Bacillus subtilis]
	Bt1G623	105-1	g1934629	131	131	1.00E-08	69	39 ((U93875) YraL [Bacillus subtilis]
	Bt1G624	914-478	g2619056	401	337	1.50E-30	58	33 ((AF027868) putative transporter [Bacillus subtilis]
498 Bt1Gc762	Bt1G625	468-1334	g2634335	550	587	4.80E-57	40	100	(Z99114) similar to macrolide glycosyltransferase [Bacillus subtilis]
499 Bt1Gc765	Bt1G626	270-40	g2633909	100	157	1.80E-11	40	.99	(Z99112) similar to hypothetical proteins [Bacillus subtilis]
499 Bt1Gc765	Bt1G627	1216-440	g133475	1164	1175	2.30E-119	89	100	RNA POLYMERASE SIGMA-G FACTOR (STAGE III SPORULATION PROTEIN G) [Bacillus subtilis]
499 Bt1Gc765	Bt1G628	1864-1377	g133289	755	719	4.90E-71	96	68	RNA POLYMERASE SIGMA-35 FACTOR PRECURSOR
500 Bt1Gc767	Bt1G629	1-1260	g1865711	592	6 <u>4</u> 6	2.70E-63	51	78	[Bacillus thuringiensis] (Y11477) endolysin [Bacteriophage Bastille]
501 Bt1Gc758	Bt1G630	1149-1	g1945050	1844	1685	2.10E-173	98		(U63928) L2 protein [Bacillus cereus]
502 Bt1Gc768	Bt1G631	1-816	g2117698	898	904	1.20E-90	63	%	aspartate carbamoyltransferase (EC 2.1.3.2) - Bacillus
503 Bt1Gc771	Bt1G632	1421-1	g2497686	69	130	6.80E-16	24	50	caldolyticus [] PLATELET-ACTIVATING FACTOR
									ACETYLHYDROLASE PRECURSOR (PAF

Table 1

Cene Id Position NCBI gi nap Score Score Prob Ident Cvrg ACETYLHYDROLASE) (LDL-SCORLATED PHOSPHOLIPASE A2) (LDL-SCORLATED PHOSPHOLIPASE A2) (LDL-PLACE) (1-1 2129188 369 369 6.00E-34 72 31 31 31 32 32 32 32 3
sat_ nap Score BlastP Prob Hent Cvrg Ident Cvrg Score Prob Ident Cvrg 369 369 6.00E-34 72 31 597 615 5.10E-60 67 47 284 300 1.80E-25 33 32 184 202 1.80E-15 91 9 217 298 2.00E-26 30 84 1171 1103 1.00E-111 65 49 417 447 3.30E-42 93 20 627 687 1.20E-67 42 62 447 473 5.70E-45 32 98 112 112 1.00E-06 27 99 408 434 7.80E-41 58 48 387 349 7.90E-32 43 99 745 660 8.80E-65 74 93 685 558 5.60E-54 72 46 596 617
BlastP BlastP- % % Score Prob Ident Cvrg 369 6.00E-34 72 31 615 5.10E-60 67 47 300 1.80E-25 33 32 202 1.80E-15 91 9 208 2.00E-26 30 84 1103 1.00E-111 65 49 447 3.30E-42 93 20 687 1.20E-67 42 62 473 5.70E-45 32 98 131 1.00E-06 27 99 434 7.80E-41 58 48 434 7.80E-41 58 48 434 7.90E-32 43 99 558 5.60E-54 72 46
BlastP BlastP- % % Score Prob Ident Cvrg 369 6.00E-34 72 31 615 5.10E-60 67 47 300 1.80E-25 33 32 202 1.80E-15 91 9 202 1.80E-15 91 9 298 2.00E-26 30 84 1103 1.00E-111 65 49 447 3.30E-42 93 20 687 1.20E-67 42 62 473 5.70E-45 32 98 131 1.00E-08 31 99 434 7.80E-155 77 71 112 1.00E-06 27 93 48 349 7.90E-32 43 99 660 8.80E-65 74 93 558 5.60E-54 72 46 617 3.20E-60 55 100 3
% % ldent Cvrg 72 31 67 47 33 32 91 9 91 9 30 84 65 49 93 20 42 62 32 98 31 99 58 48 43 99 74 93 55 100 38 76
% % Ident Cvrg 72 31 67 47 33 32 91 9 30 84 65 49 93 20 42 62 32 98 31 99 58 48 43 99 74 93 72 46 55 100 58 76
NCBI gi description ACETYLHYDROLASE) (PAF 2-ACYLHYDROLASE) (LDL-ASSOCIATED PHOSPHOLIPASE A2) (LDL- PLA(2)) (2-ACETYL-1- ALKYLGLYCEROPHOSPHOCHOLINE ESTERASE) (1- ALKYL-2-ACETYLGLYCEROPHO [Cavia porcellus] OLIGOPEPTIDE TRANSPORT ATP-BINDING PROTEIN OPPF [Bacillus subtilis] (Z99110) oligopeptide ABC transporter (ATP-binding protein) [Bacillus subtilis] (AB001488) PROBABLE TRANSPORT PROTEIN, SIMILAR PRODUCT IN MYCOBACTERIUM LEPRAE. [Bacillus subtilis] ORF IS231C [Bacillus thuringiensis] (AP000007) 306aa long hypothetical UDP-glucose 4- epimerase [Pyrococcus horikoshii] TRANSKETOLASE [Bacillus subtilis] ORF IS231C [Bacillus subtilis] ORF IS231C [Bacillus subtilis] (AF021937) catalase 1 [Arabidopsis thaliana] kinB protein - Bacillus subtilis [] (AF021937) catalase 1 [Arabidopsis thaliana] kinB protein - Bacillus megaterium] (AF065404) pX01-73 [Bacillus megaterium] (AF065404) pX01-73 [Bacillus subtilis] (Z75208) hypothetical protein [Bacillus subtilis] DEOXYRIBOSE-PHOSPHATE ALDOLASE (PHOSPHODEOXYRIBOALDOLASE) (DEOXYRIBOALDOLASE) [Bacillus subtilis] DEOXYRIBOALDOLASE) [Bacillus subtilis] (Y09323) hypothetical protein [Bacillus subtilis] (Y09323) hypothetical protein [Bacillus cereus] (U66480) YnaE [Bacillus subtilis]

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533 533	531 531 532	529 530	528	527	526 536	525	524	523	522	522	521	520	519	518	518	518	SEQ SEQ
Bt1Gc808 Bt1Gc808	Bt1Gc806 Bt1Gc806 Bt1Gc807	Bt1Gc803 Bt1Gc805	Bt1Gc802	Bt1Gc800	Bt1Gc799	Bt1Gc796	Bt1Gc797	Bt1Gc795	Bt1Gc798	Bt1Gc798	Bt1Gc794	Bt1Gc793	Bt1Gc791	Bt1Gc790	Bt1Gc790	Bt1Gc790	Contig Id
Bt1G672 Bt1G673	Bt1G669 Bt1G670 Bt1G671	Bt1G667 Bt1G668	Bt1G666	Bt1G665	Bt1G663	Bt1G662	Bt1G661	Bt1G660	Bt1G659	Bt1G658	Bt1G657	Bt1G656	Bt1G655	Bt1G654	Bt1G653	Bt1G652	Gene Id
718-191 1104-892	74-1 1331-183 1-995	1-1083 497-1366	1-1178	957-1	684-1	1-487	1440-753	1-1737	824-621	557-78	1063-112	705 - 1	1351-125	559-723	223-723	1-306	Position
g2635678 g1770028	g1146349 g1945051 g2982764	g1352313 g131027	g129021	g2127359	g2634058	g1825622	g1176993	g2501051	g2635515	g1731124	g2088525	g2313187	g137192	g2634339	g3256904	g2634340	NCBI gi
550 260	90 1 875 495	597	1524	169	918 400	97	633	778	159	658	432	359	1506	197	104	302	nap Score
562 266	106 1135 477	635 436	1481	209	924	118	645	869	168	512	450	380	1530	197	175	307	BlastP Score
2.10E-54 4.90E-23	4.50E-06 9.10E-158 2.20E-45	3.90E-62 4.80E-41	8.80E-152	5.40E-17	9.30E-93	2.30E-08	3.40E-63	6.20E-87	1.20E-12	4.20E-49	1.60E-42	8.00E-35	5.60E-157	1.00E-15	2.20E-13	2.20E-27	BlastP- Prob
59 73	92 97 36	35	73	53	<i>5</i> 75	25	53	33	43	76	35	37	72	S	26	54	% Ident
99	38 100 85	100	92	29	33 34	4	100	98	. 99	. 99	100	40	67	ω ∞	60	87	% Cvrg
aeolicus] (Z99120) yueE [Bacillus subtilis] (Z75208) hypothetical protein [Bacillus subtilis]	(L43071) ORF [Edwardsiella tarda] (U63928) L1 protein [Bacillus cereus] (AE000669) N-acetylornithine aminotransferase [Aquifex	DIPEPTIDE-BINDING PROTEIN DPPE PRECURSOR [Bacillus subtilis] PROTEIN EXPORT PROTEIN PRSA PRECURSOR	[Clostridium perfringens] SPO0B-ASSOCIATED GTP-BINDING PROTEIN	hypothetical protein 3 - Clostridium perfringens	(Z99112) similar to processing protease [Bacillus subtilis]	PROTEIN IN IDH 3'REGION [Bacillus subtilis] (U88170) No definition line found [Caenorhabditis elegans]	[Saccharomyces cerevisiae] HYPOTHETICAL 26.6 KD SENSORY TRANSDUCTION	PROBABLE ARGINYL-TRNA SYNTHETASE, CYTOPLASMIC (ARGININETRNA LIGASE) (ARGRS)	(Z99119) similar to hypothetical proteins from B. subtilis	HYPOTHETICAL 17.8 KD PROTEIN IN BLTR-SPOIIIC INTERGENIC REGION (Bacillus subtilis)	(cpdB) [Helicobacter pylori 26695] (U70619) heroin esterase [Rhodococcus sp.]	(AE000532) 2',3'-cyclic-nucleotide 2'-phosphodiesterase	EXCINUCLEASE ABC SUBUNIT C [Bacillus subtilis]	horikoshii] (Z99114) voiG 「Bacillus subtilis]	(AP000002) 272aa long hypothetical protein [Pyrococcus	[Haemophilus influenzae Rd] (Z99114) voif [Bacillus subtilis]	NCBI gi description

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SEQ NO	Contig Id	Gene Id	Position	NCBI gi	nap Score	BlastP Score	BlastP- Prob	% Ident	%	NCBI gi description
533	Bt1Gc808	Bt1G674	1220-1345	g1770029	133	133	1.20E-08	82	17	(Z75208) hypothetical protein [Bacillus subtilis]
534	Bt1Gc809	Bt1G675	1-872	g1709639	1110	1110	1.80E-112	71	71	PEPTIDASE T (AMINOTRIPEPTIDASE)
727	D+10-200	D+10676	1000 017	~ ハー・ハー・ハー・ハー・ハー・ハー・ハー・ハー・ハー・ハー・ハー・ハー・ハー・ハ	026	707	6 60E 70	3.	3	(TRIPEPTIDASE) [Bacillus subtilis]
534	Bt1Gc809	Bt1G6/6	1828-917	g477274	836	794	5.50E-79	52	100	3-methyladenine DNA glycosylase - Bacillus subtilis []
535	Bt1Gc804	Bt1G677	1605-1	g3322922	340	453	9.80E-42	27	41	(AE001237) exonuclease (sbcC) [Treponema pallidum]
536	Bt1Gc812	Bt1G678	273-1253	g1730898	1225	1112	1.10E-112	75	100	HYPOTHETICAL 36.3 KD PROTEIN IN RECQ-CMK INTERGENIC REGION [Bacillus subtilis]
537	Bt1Gc810	Bt1G679	914-1	g2506131	1046	1031	4.20E-104	69	34	ACONITATE HYDRATASE (CITRATE HYDRO-LYASE)
)					3	3	(ACONITASE) [Bacillus subtilis]
338	BUGGII	D890119	/4/-1	gysoto	1116	1016	1.70E-102	93	۰,	tlagellin B chain - Bacillus thuringiensis [Bacillus thuringiensis alestil
539	Bt1Gc814	Bt1G681	1-859	g1881339	261	108	6.60E-15	67	48	(AB001488) PROBABLE TRANSCRIPTIONAL
										REGULATOR, SIMILAR TO REGULATOR OF ANTIBIOTIC TRANSPORT COMPLEXES IN
										STREPTOMYCES HYGROSCOPICUS. [Bacillus subtilis]
539	Bt1Gc814	Bt1G682	1829-1	g1169000	262	353	5.60E-31	36	19	MICROBIAL COLLAGENASE PRECURSOR (120 KD COLLAGENASE) [Clostridium perfringens]
540	Bt1Gc813	Bt1G683	525-1092	g96807	269	359	8.70E-32	41	21	Mg2+-transporting ATPase (EC 3.6.1) mgtB - Salmonella
541	Bt1Gc817	Bt1G684	875-1	g1731017	645	562	2.10E-54	4 8	40	HYPOTHETICAL 80.1 KD PROTEIN IN SODA-COMGA
		-								INTERGENIC REGION [Bacillus subtilis]
542	Bt1Gc815	Bt1G685	1303-117	g2635180	1637	1597	4.50E-164	78	93	(Z99117) similar to protease [Bacillus subtilis]
543	Bt1Gc816	Bt1G686	1164-728	g1708641	205	242	1.70E-20	33	46	(US1115) YeaC [Bacillus subtilis]
544	Bt1Gc818	Bt1G687	102-485	g2635382	283	301	9.70E-27	4 3	99	(Z99118) similar to hypothetical proteins [Bacillus subtilis]
545	Bt1Gc819	Bt1G688	1-142	·g2633732	140	161	6.60E-12	58	22	(Z99111) similar to hypothetical proteins [Bacillus subtilis]
545	BtlGc819	Bt1G689	141-671	g2633733	643	659	1.10E-64	69	99	
343	Bt1Gc819	Bt1G690	1963-1612	g3183454	333	:- 388	5.80E-36	63	· <u>4</u>	HYPOTHETICAL 30.7 KD PROTEIN IN MCPC-KINA INTERGENIC REGION [Bacillus subtilis]
546	Bt1Gc820	Bt1G691	848-101	g2497382	606	2	9.00E-63	94	31	TRANSPOSASE FOR INSERTION SEQUENCE ELEMENT IS232 Unsertion sequence IS2321
546	Bt1Gc820	Bt1G692	565-848	g225559	447	461	1.10E-43	97	20	ORF IS231C [Bacillus thuringiensis]
547	Bt1Gc821	Bt1G693	198-1	g1434920	107	101	0.00031	39	∞	(D86346) crystal protein [Bacillus thuringiensis]
547	Bt1Gc821	Bt1G694	951-585	g2226238	452	484	3.90E-46	72	35	(Y14083) hypothetical protein [Bacillus subtilis]
548	Bt1Gc823	Bt1G695	581-234	g2522004	464	467	2.50E-44	70	99	(AB007637) function unknown [Bacillus subtilis]
549	Bt1Gc824	Bt1G696	1-1465	g1255196	1768	1787	3.30E-184	2	83	(U50744) BSMA [Bacillus stearothermophilus]
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SEQ DEQ	Contig Id	Gene Id	Position	NCBI gi	nap Score	BlastP Score	BlastP- Prob	% % Ident Cvrg	%rg	NCBI gi description
551	Bt1Gc822	Bt1G698	854-1	g2558482	1125	1130	1.40E-114	74	72	(AB008120) phosphopentomutase [Bacillus stearothermophilus]
552	Bt1Gc826	Bt1G699	1-800	g4835313	127	185	1.90E-14	33	88	(AL049863) hypothetical protein [Streptomyces coelicolor]
552	Bt1Gc826	Bt1G700	687-1455	g1644210	687	679	8.50E-67	53		(D64126) unknown [Bacillus subtilis]
553	Bt1Gc829	Bt1G702	274-1	g118797	168	182	1.50E-13	39		DNA POLYMERASE III, BETA CHAIN [Bacillus subtilis]
553	Bt1Gc829	Bt1G703	712-351	g2129134	92	156	2.20E-11	28 .	72	mutator protein mutT - Methanococcus jannaschii
) }	3)))		i ; ; i		! !	·			[Methanococcus jannaschii]
	Bt1Gc829	Bt1G704	1459-1034	g1181177	195	220	3.70E-18	30	99	(L25828) ORF 12 [Erwinia amylovora]
554	Bt1Gc830	Bt1G705	1-149	g2274866	124	143	5.30E-10	53	71	(AJ000005) orf1 [Bacillus megaterium]
554	Bt1Gc830	Bt1G706	158-972	g2274867	.997	716	1.00E-70	69	84	(AJ000005) glucose kinase [Bacillus megaterium]
555	Bt1Gc833	Bt1G707	1083-878	g3005554	. 86	113	4.00E-06	29	22	(AF047044) putative transposase [Anal
556	Bt1Gc828	Bt1G708	1929-420	g728913	1323	1234	1.30E-125	51	56	MG(2+) TRANSPORT ATPASE, P-TYPE 1 [Escherichia coli1
557	Bt1Gc834	Bt1G709	1110-1	g1730904	1129	1108	2.90E-112	59	82	HYPOTHETICAL 51.2 KD PROTEIN IN RECQ-CMK
557	Bt1Gc834	Bt1G710	1639-1129	g1644192	8 33	771	1.50E-76	94	66	(D63645) spore cortex-lytic enzyme prepeptide [Bacillus
558	Rt1Gc835	R:16711	41-598	67776757	433	121	2 30E-20	45	8	cereus] (V14084) hypothetical protein [Bacillus subtilia]
	Bt1Gc835	Bt1G712	686-1308	g2226256	344	418	3.90E-39	42		(Y14084) hypothetical protein [Bacillus subtilis]
	Bt1Gc836	Bt1G713	643-251	g629241	134	212	2.60E-17	34		lmbK protein - Streptomyces lincolnensis [Streptomyces
					v.					lincolnensis]
559	Bt1Gc836	Bt1G714	1162-1	g3915419	47	124	5.50E-08	36	56	HYPOTHETICAL 17.9 KD PROTEIN IN PHOB-GROES INTERGENIC REGION [Bacillus subtilis]
560	Bt1Gc839	Bt1G715	256-1	g2634117	264	287	2.90E-25	60	27	(Z99113) tRNA isopentenylpyrophosphate transferase
560	Bt1Gc839	Bt1G716	1152-751	g2619050	277	319	1.20E-28	52	45	[Bacillus subtilis] (AF027868) YocH [Bacillus subtilis]
561	Bt1Gc838	Bt1G717	1246-468	g586817	998	999	1.00E-100	77	84	HYPOTHETICAL 34.0 KD PROTEIN IN COTF-TETB
\$60	Rt1Gc840	Rt1G718	1106-203	03355675	261	363	2 60E-33	3		INTERGENIC REGION [Bacillus subtilis]
	0.000		1100-200	80000000	. 101	Ü	4.00E-00	36	. 0	(ALV31124) hypometical protein 3C1C2.ve [surficilityees coelicolor]
563	Bt1Gc841	Bt1G719	488-710	g4874247	80	138	3.30E-09	34	29	(AF041061) beta-lactamase-like protein [Mycobacterium smeamatis]
	Bt1Gc842	Bt1G720	198-1391	g585250	131	218	6.00E-18	28		
565	Bt1Gc843	Bt1G721	1451-315	g1731316	1136	1157	1.90E-117	55	100	HYPOTHETICAL 41.6 KD PROTEIN IN IDH-DEOR

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579 Bt1Gc865 Bt1G744 1507-326	Bt1Gc863 Bt1G743	578 Bt1Gc863 Bt1G742 354-1	577 Bt1Gc858 Bt1G741 1239-1567	577 Bt1Gc858 Bt1G740 871-1		576 Bt1Gc855 Bt1G739 1174-1537	576 Bt1Gc855 Bt1G738 972-607		576 Bt1Gc855 Bt1G737 393-1	575 Bt1Gc853 Bt1G735 1879-1			574 Bt1Gc857 Bt1G734 1309-1477		574 Bt1Gc857 Bt1G733 . 10-717	Bt1Gc852 Bt1G732	572 Bt1Gc854 Bt1G731 1134-1562	572 Bt1Gc854 Bt1G730 766-503	572 Bt1Gc854 Bt1G729 1-424	-	571 Bt1Gc845 Bt1G728 1-1178		570 Bt1Gc832 Bt1G727 1152-619	Bt1Gc847 Bt1G726	568 Bt1Gc844 Bt1G725 384-1333	568 Bt1Gc844 Bt1G724 1-381		Bt1Gc850 Bt1G723	566 Bt1Gc846 Bt1G722 1-1371		D Contig Id Gene Id Position	SEO
326 g1684651	-	1 g2340007	567 g3258280			537 g139993			1 g4033506		•		477 g1881344	•	17 g2636560		-	03	4 g2633836	•	78 g1731015	1	619 g1176958			1 g132678			71 62619033		NCBI gi	
638 349		345 345	112 139	803 792		206 240	594 594		582 583	1890 1770			103 128		390 403		537 530	260 263	327 339		1139 955		132 8		1153 990	541 486	-		1606 1624	: :	nap Score Score	991
9 9.00E-55		5 2.10E-31	9 1.40E-09	2 9.00E-79		0 2.80E-20	4 8.60E-58		3 1.30E-56	0 2.10E-182			8 1.90E-07		3 1.50E-37		0 5.20E-51	3 1.00E-22	9 9.10E-31	. •	5 4.80E-96		80 3.60E-08		0 9.40E-100	6 2.40E-46			4 6 20E-167		e Prob	
33 100	58	54 45	39 63	54 46) 45 68	98 99		88 45	94			40 12		7 37 100	36 1	74 29	55 99	45 67		56 91		3 29 99	32	-	84 86	.~	53 (60 5		% % Ident Cvrg	
_				_	INTERGENIC REGION (ORFS) [Bacillus subtilis]		9 spo0F gene [Bacillus thuringiensis]	stearothermophilus]	5 FRUCTOSE-BISPHOSPHATE ALDOLASE [Bacillus		MELILOTI. [Bacillus subtilis]	CATABOLISM (MOCR) GENE OF RHIZOBIUM		[Bacillus subtilis]		_		_	_	INTERGENIC REGION [Bacillus subtilis]		INTERGENIC REGION [Bacillus subtilis]					(AUTOLYSIN) [Bacillus sp.]			INTERGENIC REGION [Bacillus subtilis]	NCBI gi description	

Table 1

INTERGENIC REGION [Escherichia conj								.•		
PRECURSOR (CELL WALL HYDROLASE) (AUTOLYSIN) [Bacillus subtilis] HYPOTHETICAL 25.3 KD PROTEIN IN NFO-FRUA	100	30	9.00E-24	273	244	g465600	475-1152	Bt1G762	Bt1Gc874	590
N-ACETYLMURAMOYL-L-ALANINE AMIDASE XYLA	42	55	9.80E-18	216	303	g731171	389-1	Bt1G761	Bt1Gc873	589
(AE001237) exonuclease, putative [Treponema pallidum]	77	24	1.70E-18	227	100	g3322921	2128-1249	Bt1G760	Bt1Gc872	588
[riasmodium raiciparum] (AE001237) exonuclease (sbcC) [Treponema pallidum]	38		5.40E-26	306	230	g3322922	1243-1	Bt1G759	Bt1Gc872	588
nver stage antigen LSA-1 - Flasmodium tatciparum	20	17	6.90E-11	100	09	807/039	1-1007	,007,0110	שנוטנס/2	000
INTERGENIC REGION [Bacillus subtilis]	3	1	0 00E 11	105	ŝ	0507050	2061 1	B+1C749	B+1C-872	488
HYPOTHETICAL 23.6 KD PROTEIN IN QCRC-DAPB	43	74	4.90E-32	351	333	g1176701	1414-1138	Bt1G757	Bt1Gc871	587
horikoshii]										
(AP000006) 197aa long hypothetical protein [Pyrococcus	99	37	7.80E-25	283	191	g3257896	435-1100	Bt1G756	Bt1Gc871	587
(M29945) ORF 2; putative [Yersinia enterocolitica]	94	54	2.70E-31	344	334	g1197032	1-353	Bt1G755	Bt1Gc871	. 587
(AJ002571) YkoD [Bacillus subtilis]	76	33	2.30E-50	524	452	g2632042	1-1141	Bt1G754	Bt1Gc862	. 586
[Bacillus subtilis]								٠		
ENZYME II, ABC COMPONENT) (EII-GLC / EIII-GLC)							·			
IIABC COMPONENT) (PHOSPHOTRANSFERASE							,			
COMPONENT (EIIABC-GLC) (GLUCOSE-PERMEASE						•		`		
PTS SYSTEM, GLUCOSE-SPECIFIC IIABC	46	73	3.50E-109	1079	1182	g131493	1-968	Bt1G753	Bt1Gc869	585
(Z99104) ybcF [Bacillus subtilis]	71	52	3.90E-23	267	289	g2632453	1486-1108	Bt1G752	Bt1Gc868	584
ACETYL ESTERASE [Escherichia coli]	49	26	3.30E-11	159	57	g2506820	495-1	Bt1G751	Bt1Gc868	584
[Pseudomonas oleovorans]					-					
(MEDIUM-CHAIN ACYL-COA SYNTHETASE)										
MEDIUM-CHAIN-FATTY-ACIDCOA LIGASE	8	42	1.90E-108	1072	1013	g416605	1437-1	Bt1G750	Bt1Gc861	583
HEMA INTERGENIC REGION (ORFX) [Bacillus subtilis]										
HYPOTHETICAL GTP-BINDING PROTEIN IN LONA-	43	45	1.60E-15	195	158	g586754	1490-1239	Bt1G749	Bt1Gc867	582
(AJ002571) YkhA [Bacillus subtilis]	99	51	7.70E-34	368	319	g2632036	780-1190	Bt1G748	Bt1Gc867	582
(AJ002571) YkmA [Bacillus subtilis]	99	50	3.30E-33	362	342	g2632035	315-756	Bt1G747	Bt1Gc867	582
embrilie]		i				9				,
[Synechocystis sp.] (Z99114) similar to macrolide olycosyltransferase [Bacillus	.	42	4.60E-59	606	570	g2634335	366-1224	Bt1G746	Bt1Gc866	581
(ASPARAGINETRNA LIGASE) (ASNRS)										
PROBABLE ASPARAGINYL-TRNA SYNTHETASE	65	63	1.10E-112	1112	1120	g1711644	1269-263	Bt1G745	Bt1Gc860	580
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NCBI gi description	%Cvrg	% % Ident Cvrg	BlastP- Prob	BlastP Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	8 B 8
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HYPOTHETICAL 36.3 KD LIPOPROTEIN PRECURSOR	23	37	2.20E-10	152	136	g732409	1250-1012	Bt1G786	Bt1Gc890	605
PROBABLE ABC TRANSPORTER PERMEASE	100	61	7.10E-63	642	864	g732410	1006-179	Bt1G785	Bt1Gc890	605
INSERTION SEQUENCE IS232 PUTATIVE ATP-	32	99	1.70E-38	412	412	g2497392	880-641	Bt1G784	Bt1Gc892	604
(D86418) YfnH [Bacillus subtilis]	65	54	2.10E-47	496	499	g2116766	497-1	Bt1G783	1 Bt1Gc892	604
(AF047044) putative transposase [Anabaena PCC7120]	28	29	2.60E-09	142	79	g3005554	1-270	Bt1G782	Bt1Gc891	603
(D86380) Alkaline D-peptidase [Bacillus cereus]	100	25	2.20E-27		152	g1741967	151-1295	Bt1G781	Bt1Gc889	602
horikoshii]						(
(AP000001) 289aa long hypothetical protein [Pyrococcus	100	29	1.20E-30	338	241	g3256528	267-1121	Bt1G780	Bt1Gc889	602
ATP SYNTHASE GAMMA CHAIN [Bacillus megaterium]	68	53	4.00E-53	550	520	g114636	587-1	Bt1G779	Bt1Gc888	601
	5	č	1.001-00			82022102	110000	51.0	, 201000	9
		3	2 90E 6	603	603	~2625100	1155 610	D+1C770		2
	83	66	1.50E-51	535	619	g2635190	541-1	Bt1G777	Bt1Gc885	. 600
[Aquifex aeolicus]										
(AE000682) transcriptional regulator (TetR/AcrR family)	73	31	1.50E-12	167	84	g2982980	752-1	Bt1G776	Bt1Gc884	599
(Z75208) hypothetical protein [Bacillus subtilis]	10	69	4.30E-08	132	116	g1770012	1-106	Bt1G775	Bt1Gc884	599
aeolicus]										
(AE000675) NADH dehydrogenase (ubiquinone) [Aquifex	45	25	3.20E-10	150	104	g2982870	586-1003	Bt1G774	Bt1Gc881	598
(U87792) unknown [Bacillus subtilis]	99	73	8.60E-58	594	566	g1842437	921-446	Bt1G773	7 Bt1Gc883	597
(Z99112) ymfL [Bacillus subtilis]	99	54	2.80E-20	· 240	223	g2634062	446-172	Bt1G772	7 Bt1Gc883	597
(Y08953) cytidylate kinase-like protein [Bacillus cereus]	51	98	9.10E-47	490	487	g1945144	1340-1037	Bt1G771	Bt1Gc882	596
305 KIBUSUMAL FRUTEIN ST HUMULUG [Bacillus	0	90	4.80E-112	1100	1102	810771CB	1-660	P110//0) P(1/0/2007	. 390
(AFUIS//S) YOUC [Bacillus subtilis]	10	۲ ۲	8.10E-46	481	1102	g2415385	103-/08	B110759		600
(AL049485) hypothetical protein [Streptomyces coelicolor]	.73	22	5.30E-19	228	155	g4539178	216-788	Bt10768		794
(AF008220) transporter [Bacillus subtilis]	73	50	2.30E-48	505	468	g2293177	2285-1737	Bt1G767		. 593
(D14399) hypothetical protein [Bacillus subtilis]	.100	41	7.70E-50	519	655	g709994	1744-512	Bt1G766		593
[Bacillus subtilis]										
1	91	38	1.30E-68	696	777	g1731300	1744-1	Bt1G765	Bt1Gc875	593
						•		. :		
subtilis] (D88802) B. subtilis alkaline phosphatase IIIA; P19405	97	2	1.90E-140	1374	1423	g1945090	1-1341	Bt1G764	Bt1Gc877	592
STAGE IV SPORULATION PROTEIN A [Bacillus	100	88	1.60E-216	2092	2244	g464794	1705-234	Bt1G763	Bt1Gc876	591
NCBI gi description	Cyrg	Ident	Prob	Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	8 8
	%	%	BlastP-	BlastP	aat			1		SEQ

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NODULATION PROTEIN NOLG PRECURSOR	39	. 29	3.40E-11	157	132	g128506	825-503	Bt1G810	Bt1Gc910	624
(AE001125) acriflavine resistance protein (acrB) [Borrelia	27	•	1.70E-13	189	183	g2688027	822-1	Bt1G809	Bt1Gc910	624
collagen adhesin - Staphylococcus aureus []	16	27	2.40E-15	192	47	g282243	1-1676	Bt1G808	Bt1Gc913	623
(Z99120) yunE [Bacillus subtilis]	91		6.10E-41	435	676	g2635735	895-1625	Bt1G807	Bt1Gc912	622
(AJ002571) YkaB [Bacillus subtilis]	77	77	1.10E-57	593	958	g2632005	1-760	Bt1G806	Bt1Gc912	622
(X92868) NAD alcohol dehydrogenase [Bacillus subtilis]	52		4.00E-37	399	269	g2108273	1405-1	Bt1G805	Bt1Gc911	621
(X99724) EA1 [Bacillus anthracis]	9		3.40E-10	157	118	g1769919	229-1	Bt1G804	Bt1Gc911	621
INTERGENIC REGION [Bacillus subtilis]							ļ.)))	<u>;</u>
(ORF 2) [Clostridium perfringens] HYPOTHETICAL 47.6 KD PROTEIN IN EPR-GALK	58	42	5.70E-38	407	450	g732331	1412-649	Bt1G803	Bt1Gc907	620
HYPOTHETICAL 55.7 KD PROTEIN IN PLC 5'REGION	90	62	6.10E-144	1407	1384	g549806	1318-1	Bt1G802	Bt1Gc909	619
LYTB PROTEIN HOMOLOG [Bacillus subtilis]	. 25		4.90E-32	351	344	g1731004	1001-1244	Bt1G801	Bt1Gc905	618
(L48554) vrrA gene product [Bacillus anthracis]	100		6.70E-40	235	1026	g1162917	177-908	Bt1G800	Bt1Gc905	618
(Z99121) yvbH [Bacillus subtilis]	91	60	1.70E-52	544	613	g2635899	1306-747	Bt1G799	Bt1Gc904	617
(ALOST 101) phante (wo-component regulator	6	Ų	1.001-24	707	2.23	80007770	101-1	D110770	P(10070)	9
	6		1 005 7	ູ	260	~222A700	161 1	B+1.C709	B+1C-003	۲۱۷ د
(AF025433) tyrosine/dopa decarboxylase [Papaver	89	27	3.70E-50	522	436	g3282523	1344-46	Bt1G797	Bt1Gc902	615
(Z82044) hypothetical 54.4 kd protein [Bacillus subtilis]	62		5.20E-51	530	480	g1673402	748-1623	Bt1G796	Bt1Gc899	614
subtilis]						•				
(Z99107) similar to acriflavin resistance protein [Bacillus	66	37	2.30E-96	958	1127	g2632985	1-2094	Bt1G795	Bt1Gc898	613
STAGE V SPORULATION PROTEIN S [Bacillus subtilis]	. 99	_	2.30E-09	137	318	g1174408	614-357	Bt1G794	Bt1Gc901	612
INTERGENIC REGION [Bacillus subtilis]										
HYPOTHETICAL 10.9 KD PROTEIN IN PHRC-GDH	53	39	3.00E-07	117	102	g3183248	1-151	Bt1G793	Bt1Gc900	611
HYPOTHETICAL 58.3 KD PROTEIN IN GLPD-CSPB	83	4	1.70E-75	761	954	g3915990	1549-241	Bt1G792	Bt1Gc896	610
[Bacillus subtilis]			•							
(AB001488) PROBABLE REPRESSOR PROTEIN.	98	35	4.90E-16	200	136	g1881293	1-404	Bt1G791	Bt1Gc897	609
denendent alpha subunit) [Bacillus subtilis]						(
(Z99108) acetoin dehydrogenase E1 component (TPP-	94		1.90E-133	1308	1329	g2633130	1-946	Bt1G790	Bt1Gc895	608
(Z75208) autolysin response regulator [Bacillus subtilis]	97	28	2.30E-18	222	189	g1770002	1396-2083	Bt1G789	Bt1Gc893	607
(AB000617) Ycel [Bacillus subtilis]	98	69	4.00E-147	1437	1445	g2415746	1-1177	Bt1G788	Bt1Gc893	607
(Z99110) similar to hypothetical proteins [Bacillus subtilis]	100	36	1.60E-43	435	509	g2633695	44-972	Bt1G787	Bt1Gc894	606
(ORFK) [Bacillus subtilis]					٠.					
NCBI gi description	Cyrg	Ident Cyrg	Prob	Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	8 8
	8	\$	BlastP-	BlastP	aat:		-	,	I	SEQ
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Table 1

634	633	633	632	632	631	631	630	629	628	627	627		626	•	626	626	625	625	NO TEQ SEQ
Bt1Gc926	Bt1Gc928	Bt1Gc928	Bt1Gc924	Bt1Gc924	Bt1Gc923	Bt1Gc923	Bt1Gc922	Bt1Gc921	Bt1Gc920	Bt1Gc917	Bt1Gc917		Bt1Gc919		Bt1Gc919	Bt1Gc919	Bt1Gc915	Bt1Gc915	Contig Id
Bt1G827	Bt1G826	Bt1G825	Bt1G824	Bt1G823	Bt1G822	Bt1G821	Bt1G820	Bt1G819	Bt1G818	Bt1G817	Bt1G816		Bt1G815		Bt1G814	Bt1G813	Bt1G812	Bt1G811	Gene Id
1523-1093	680-549	469-115	1246-1901	1-1227	1813-554	554-1	684-1361	1-1657	1444-866	1801-334	1-243		1942-1615		1611-262	177-1	1-720	1-762	Position
g2224769	g132902	g585905	g417313	g417328	g2633730	g2633731	g1684650	g2621914	g2313950	g4584148	g4584149		g417313		g417327	g134774	g3127079	g3114663	NCBI gi
394	204	337	764	1188	1212	550	380	217	119	2453	274	•	475	•	1279	213	412	428	nap Score
318	204	365	639	1154	1288	524	415	436	136	2175	280		394		1300	168	329	312	BlastP Score
1.50E-28	1.80E-16	1.60E-33	1.50E-62	3.90E-117	2.50E-131	2.30E-50	8.00E-39	1.20E-40	5.70E-18	2.50E-225	1.60E-24		1.30E-36		1.30E-132	5.00E-12	1.00E-29	6.60E-28	BlastP- Prob
3 55	5 91	3 59	2 68	7 57	60) 53	39) 29	3 25	5 97	75		85		2 55	. 69	37	37	% Ident
. 69	9.	95	67	83	100	78	47	·52	100	50	26		34	. :	100		86	. 90	%Cvrg
stearothermophilus] (Z97025) product similar to B. subtilis YhcN hypothetical	C5) (RNASE P) [Bacillus subtilis] 50S RIBOSOMAL PROTEIN L34 [Bacillus	TRANSFERASE (UDP-MURNAC-PENTAPEPTIDE PHOSPHOTRANSFERASE) [Bacillus subtilis] RIBONUCLEASE P PROTEIN COMPONENT (PROTEIN	N-ACETYLMURAMYL-TRIPEPTIDE SYNTHETASE) [Bacillus subtilis] PHOSPHO-N-ACETYLMURAMOYL-PENTAPEPTIDE-	[Bacillus subtilis] UDP-N-ACETYLMURAMOYLALANYL-D-	(Z99111) similar to ribulose-bisphosphate carboxylase	(Z99111) similar to hypothetical proteins [Bacillus subtilis]	[Methanobacterium thermoautotrophicum] (Z82987) unknown [Bacillus subtilis]	pylori 26695] (AE000859) sensory transduction histidine kinase	(AE000593) osmoprotection protein (proV) [Helicobacter	(AJ010111) pyruvate carboxylase [Bacillus cereus]	(AJ010111) cytochrome aa3 controlling protein [Bacillus	TRANSFERASE (UDP-MURNAC-PENTAPEPTIDE PHOSPHOTRANSFERASE) [Bacillus subtilis]	GLUTAMIC ACID ADDING ENZYME) [Bacillus subtilis] PHOSPHO-N-ACETYLMURAMOYL-PENTAPEPTIDE-	GLUTAMATE LIGASE (UDP-N-ACETYLMURANOYL-L-ALANYL-D-GLUTAMATE SYNTHETASE) (D-			[Pseudomonas stutzeri] (AF061070) PtxC [Pseudomonas stutzeri]	[Rhizobium meliloti] (AF061267) inner membrane component HtxC	NCBI gi description

	59 84	2.30E-167	1628 2	1609	g2116756	1587-1	Bt1G848	Bt1Gc914	649
uperculosisj 5 PROBABLE 1,4-DIHYDROXY-2-NAPHTHOATE 6 OCTAPRENYLTRANSFERASE (DHNA-	60 75	5.90E-59	605	720	g732327	701-1	Bt1G847	Bt1Gc941	648
	32 16	3.20E-05	108	8	g2114014	1282-1026	Bt1G846	Bt1Gc938	647
_	28 65	2.10E-24	279	231	g1805447	738-1	Bt1G845	Bt1Gc937	646
2 PEPTIDASE T (AMINOTRIPEPTIDASE) (TRIPEPTIDASE) [Bacillus subtilis]	63 32	2.20E-43 (458	4	g1709639	389-1	Bt1G844	Bt1Gc939	645
				. 110	g1789981	1-169	Bt1G843	Bt1Gc936	4
	_	•		1676	g2633857	1-1193	Bt1G842	Bt1Gc933	643
		1.50E-23		269	g1149666	1487-1774	Bt1G841	Bt1Gc935	642
	28 . 85	3.30E-33		288	g3426013	704-1774	Bt1G840	Bt1Gc935	643 2
jannaschii] 0 (U34772) ORF375 [Dichelobacter nodosus]	35 90	1.40E-41	441	320	g1002992	710-1774	Bt1G839	Bt1Gc935	642
	26 50	1.10E-06	114	85	g2495742	427-750	Bt1G838	Bt1Gc935	6 <u>4</u> 2
PROTEIN 15) (VEG15) [Bacillus subtilis]									
NUCLEOTIDYLTRANSFERASE (POLYNUCLEOTIDE PHOSPHORYLASE) (PNPASE) (VEGETATIVE									
	83 90	3,40E-276	2655 3	2674	g1709698	Jan-09	Bt1G837	Bt1Gc916	24
7 (AF027868) PEP synthase [Bacillus subtilis]	66 37	1.40E-103	1026 1	1064	g2619033	1833-874	Bt1G836	Bt1Gc932	640
	39 29	1.30E-15	198	187	g548899	1441-1720	Bt1G835	Bt1Gc934	639
	40 40	1.00E-24 '	282	240	g2896708	418-1	Bt1G834	Bt1Gc934	639
_	49 \ 100	1.90E-78 '		753	g2632008	1142-210	Bt1G833	Bt1Gc930	638
_	49 99	9.80E-18	216	196	g1161061	1142-876	Bt1G832	Bt1Gc930	638
INTERC					•				
<pre>subtilis] 3 HYPOTHETICAL OXIDOREDUCTASE IN GNTR-GGT</pre>	46 23	1.80E-15	198	156	g1176282	1033-1274	Bt1G831	Bt1Gc931	637
UBIQUINONE OXIDOREDUCTASE CHAIN 5) [Bacillus									
	69 36	1.40E-50	526	630	g1171678	1001-1548	Bt1G830	Bt1Gc929	636
	77 99	6.10E-41		492	g2632455	31-402	Bt1G829	Bt1Gc929	636
	27 54	0.0018	101	169	g3322295	1521-955	Bt1G828	Bt1Gc927	635
-	nt Cvrg	Prob Ident	Score	Score	o d			ď	S S
NCBI gi description		BlastP- %		•	NCBI gi	Position	Gene Id	Contig Id	SEQ

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SEQ ID NO 649 651 652	Contig Id Bt1Gc914 Bt1Gc942 Bt1Gc943 Bt1Gc947	Gene Id Bt1G849 Bt1G850 Bt1G851 Bt1G852	Position 2231-1580 201-889 1-465 235-1010	NCBI gi g1730910 g2833393 g1934657 g1763710	88t_nap Score 353 560 153 263	BlastP Score 392 513 201 293	BlastP- Prob 2.20E-36 3.30E-49 8.20E-16 6.80E-26	% Ident C 41 47 43 31		NCBI gi description HYPOTHETICAL 23.1 KD PROTEIN IN BSAA-ILVD INTERGENIC REGION [Bacillus subtilis] SAPB PROTEIN [Bacillus subtilis] (U93876) hypothetical protein YrdP [Bacillus subtilis] (Z83337) highly similar to phosphotransferase system
652	2 Bt1Gc947	Bt1G853	552-1549	g2621392	75	175	2.20E-13		28	28 99 (AE000818) conserved protein [Methanobacterium thermoautotrophicum]
653 653	Bt1Gc945 Bt1Gc945	Bt1G854 Bt1G855	456-1 1210-453	g1934810 g1934809	358 554	361 590	4.20E-33 2.30E-57		45	4 4
654	1 Bt1Gc944	Bt1G856	1552-1	g3915461	1425	1419	3.30E-145		50	50 89
655		Bt1G857	2158-1397	g3059028	311	301	2.80E-26	•	32	32 50
656	5 Bt1Gc948	Bt1G858	432-1	g2811132	402	423	1.10E-39	9	56	56 26
656	5 Bt1Gc948	Bt1G859	1281-1	g4894306	455	539	5.80E-52	Ň	49	
657	7 Bt1Gc951	Bt1G860	1797-1221	g4914333	418	398	5.10E-37	2	46	46 80
658	3 Bt1Gc946	Bt1G861	880-34	g1171920	801	616	4.00E-60	Õ	59	
658	BtlGc946	Bt1G862	1891-839	g1805370	1037	968	2.00E-97	<u> </u>	60	60 84
. 039	/ BITUCY33	B110803	439-1209	8112022	1294	11/9	8.80E-120		98	
660		Bt1G864	1710-1785	g2497382	109	125	3.40E-07	-)7 96	96 .6
661	Bt1Gc954 Bt1Gc954	Bt1G865	918-208	g2634845 g1731058	116 495	122 459	9.00E-08 1.70E-43	டங்க	43 &	
.661	1 Bt1Gc954	Bt1G867	842-1098	g586811	135	162	5.20E-12	į,	3 8	38 97
662	2 Bt1Gc957	Bt1G868	Jan-16	g1731087	171	260	2.10E-22	<i>i</i> 3	30 .	30 . 59
663	3 Bt1Gc956	Bt1G869	66-1889	g1881242	2068	1683	3.40E-173	- 73	66	66 100
664	4 Bt1Gc958	Bt1G870	1386-708	g2621255	150	191	4.40E-15	i.n	24	24 100

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(299100) similar to nypometicai proteins [Bacillus subulis]	100	94	4.YUE-33	500	J Q #	82032040	007-200	DK80119	6/69/11	0/0
			2.40E-156	1524	1600	g421457	41-1336	Bt1G889		. 675
	· 30	. 65	6.40E-14	180	180	g732387	1210-1347	Bt1G888	Bt1Gc972	674
	3 99	7 43	2.90E-27	306	279	g732386	675-1115	Bt1G887	Bt1Gc972	674
				į	·į	9				
,OdAH			1.10E-39	423	423	g1730951	1-426	Bt1G886		674
subtilis] (Z99114) yoaS [Bacillus subtilis]	99	5 74	2.40E-55	571	618	g2634266	1806-1327	Bt1G885	Bt1Gc970	673
[Streptococcus pneumoniae] (Z99114) similar to transcriptional regulator [Bacillus	8 1	7 79	2.20E-27	307	296	g2634267	1313-1076	Bt1G884	Bt1Gc970	673
	97	30	3.30E-10	145	80	g2126635	1-398	Bt1G883	Bt1Gc970	673
(LIMIT DEXTRINASE) (ISOMALTASE) (DEXTRIN 6-ALPHA-D-GLUCANOHYDROLASE) [Bacillus cereus]										
_	7 92	97	3.70E-279	2683	2733	g129005	721-2268	Bt1G882	Bt1Gc966	672
_	42	80	4.90E-55	568	604	g3688811	458-1	Bt1G881	Bt1Gc966	672
(D83967) TreC [Bacillus subtilis]			3.60E-123	1211	1161	g2626830	105-1084	Bt1G880	Bt1Gc967	671
INTERGENIC REGION [Escherichia coli]										
	- 88	5 54	1. 8 0E-66	676	625	g418534	880-1	Bt1G879	Bt1Gc969	670
	§ 42	38	3.00E-39	419	385	g1075694	1246-1917	Bt1G878	Bt1Gc965	669
INTERGENIC REGION [Bacillus subtilis] HYPOTHETICAL PROTEIN (ORF3) [Bacillus cereus]	. 89	95	6.00E-121	1190	1327	g3183483	1-836	Bt1G877		669
	100	59	3.50E-61	626	625	g732345	340-939	Bt1G876	Bt1Gc964	668
ambotaciens] (Z75208) hypothetical protein [Bacillus subtilis]	.99	23	1.00E-15	197	164	g1770039	1136-552	Bt1G875	Bt1Gc963	667
	3 E	38	1.00E-08	131	103	g417332	88-1361	Bt1G874	Bt1Gc963	667
	71	56	1.20E-49	517	488	g135583	987-486	Bt1G873	Bt1Gc960	666
(Z99110) similar to hypothetical proteins [Bacillus subtilis] (AE000681) hypothetical protein [Aquifex aeolicus]	99	22 65	1.60E-08 1.10E-09	129 140	129 86	g2633504 g2982976	1-111 428-906	Bt1G871 Bt1G872	Bt1Gc961 Bt1Gc961	665
thermoautotrophicum]										
NCBI gi description	% % Ident Cvrg	% Ident	BlastP- Prob	nap Score Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	NO E
			Tadel							

Table 1

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	703 1	. 703 I	-	702	701	700 I	699 I	699 I	698 I	697	697	696	696		695	695				692	692	691			> -	₽
	Bt1Gc1003	Bt1Gc1003	Bt1Gc999	Bt1Gc999	Bt1Gc996	Bt1Gc1002	Bt1Gc1000	Bt1Gc1000	Bt1Gc1001	Bt1Gc998	Bt1Gc998	Bt1Gc997	Bt1Gc997		Bt1Gc995	Bt1Gc995	Bt1Gc991	Bt1Gc993		Bt1Gc990	Bt1Gc990	Bt1Gc992	Bt1Gc992	P(100)0)	D+1C-2000	Contig Id
	Bt1G932	Bt1G931	Bt1G930	Bt1G929	Bt1G928	Bt1G927	Bt1G926	Bt1G925	Bt1G924	Bt1G923	Bt1G922	Bt1G921	Bt1G920		Bt1G919	Bt1G918	Bt1G917	Bt1G916		Bt1G915	Bt1G914	Bt1G913	Bt1G912	1160119	510011	Gene Id
	1357-787	739-1	662-1705	1-514	1077-415	1-1405	995-1243	1-650	723-151	1943-1032	992-1	406-1053	1-214		1278-1509	52-945	1228-211	1-1020		1984-2267	1231-1602	1-1128	1-1128	1002-1040	1000 1545	Position
	g1731037	g1731038	g72992	g1842440	g2822199	g3329623	g1730936	g3183575	g2983883	g1945050	g1945051	g2635182	g2635183		g135583	g118407	g1665720	g2415733	(<u>e</u> 2145816	g3915567	g79972	g1694898	82000102	27633133	INCDI SI
	414	.204	1470	406	490	52	127	541	170	1074	1399	559	193		195	452	1623	210		117	496	194	597	. 2	Score	nap
	366	183	1401	443	534	140	127	509	225		904	580	162		221	511	_			4	458	299	635	04	ğ	חלים לים לים לים לים לים לים לים לים לים
	1.20E-33	3.10E-14	2.60E-143	8.70E-42	2.00E-51	4.80E-09	2.60E-08	8.80E-49	1.10E-18	2.00E-90	1.20E-104	2.60E-56	2.20E-11		2.90E-18	5.40E-49	4.70E-169	2.10E-22		4.20E-10	2.20E-43	2.70E-25	1.70E-61	1.000-00	1 500 36	J
	£	1 52		2 53	1 45	25	31	47	30	72	- 84	50	58		3 56	37		24		28	73	26		Ų	C.3	
	99	38	100	41	100	43	99	89	100	68	86	100	20		33	100	93	94		99	99	34	37	ą	7)
PEPTIDASE I) [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis] SIGNAL PEPTIDASE I W (SPASE I) (LEADER	HYPOTHETICAL 28.3 KD PROTEIN IN COMGG-SINR	recE protein - Bacillus subtilis []	crista] (U87792) CinA [Bacillus subtilis]	elegans (U96166) ATP-binding cassette protein [Streptococcus	[Bacillus subtilis] (AF078790) No definition line found [Caenorhabditis	INTERGENIC REGION (ORF19) [Bacillus subtilis] HYPOTHETICAL 10.2 KD PROTEIN IN ILVA 3'REGION	HYPOTHETICAL 28.1 KD PROTEIN IN PHOD-PCP	(AE000743) O-methyltransferase [Aquifex aeolicus]	(U63928) L2 protein [Bacillus cereus]	[Bacillus subtilis] (U63928) L1 protein [Bacillus cereus]	(Z99117) similar to caffeoyl-CoA O-methyltransferase	(Z99117) similar to folate metabolism [Bacillus subtilis]	subtilis]	ALANYLALANINE SYNTHETASE) [Escherichia coli] TRANSCRIPTIONAL ACTIVATOR TENA [Bacillus	D-ALANINED-ALANINE LIGASE B (D-	(D17312) diarrheal toxin [Bacillus cereus]	(AB000617) RapJ [Bacillus subtilis]	leprae [Mycobacterium leprae]	INTERGENIC REGION [Bacillus subtilis] bacitracin resistance protein homolog bacA - Mycobacterium	transposon In917 [Transposon In917] HYPOTHETICAL 14.8 KD PROTEIN IN HOM-MRGA	transposase tripA - Enterococcus faecalis plasmid pAD2	(Y09450) transposase [Pseudomonas putida]	(dihydrolipoamide acetyltransferase) [Bacillus subtilis]	(700100)	incon graceri peron

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SEQ ID Contig Id NO	Gene Id	Position	NCBI gi	nap Score	BlastP Score	BlastP- Prob	% % Ident Cvrg	% grv %	NCBI gi description
704 Bt1Gc1005	Bt1G933	1772-225	g2266421	1738	1787	3.30E-184	65	100	(Y13917) yngK [Bacillus subtilis]
	Bt1G934	772-1	g416901	1007	866	1.30E-86	77		ASPARTATE-SEMIALDEHYDE DEHYDROGENASE
705 B+1C-1006	B+1G03%	1421_022	c/17921	785	793	0 00E-70	74	100	(ASA DEHYDROGENASE) [Bacillus subtilis]
	Bt1G936	1691-1151	g1731048	607	609	2.20E-59	6 5 :	51	
								٠	INTERGENIC REGION [Bacillus subtilis]
707 Bt1Gc1008	Bt1G937	690-1	g2116761	613	589	2.90E-57	53	57	(D86418) YfnC [Bacillus subtilis]
707 Bt1Gc1008	Bt1G938	1272-893	g3868866	82	113	1.10E-05	23	22	(AB014075) GTPase [Clostridium histolyticum]
708 Bt1Gc1004	Bt1G939	167-1478	g2688567	642	614	6.60E-60	37	100	(AE001165) Na+/H+ antiporter (nhaC-1) [Borrelia
709 Bt1Gc1009	Br1G940	1435-323	03915568	1350	1360	6 50E-140	71	<u> </u>	burgdorferi] HYPOTHETICAL 39.4 KD OXIDOREDLICTASE IN
			•						HOM-MRGA INTERGENIC REGION [Bacillus subtilis]
710 Bt1Gc1010	Bt1G941	1821-1563	g2632030	224	234	1.20E-19	53	82	(AJ002571) YkkD [Bacillus subtilis]
711 BtlGc1011	Bt1G942	1245-1	g3876615	. 805	801	1.00E-79	43	48	(274031) Similarity to Yeast D-lactate dehydrogenase
								•	from this gene; cDNA EST EMBL:C12916 comes from this
									gene; cDNA EST EMBL:C10532 comes from this gene; cDNA EST EMBL:C10979 comes f []
712 Bt1Gc1012	Bt1G943	1-646	g2274944	529	479	1.30E-45	46	54	(AJ000346) NapC protein [Enterococcus hirae]
713 Bt1Gc1013	Bt1G944	347-1	g225559	560	572	1.90E-55	98	24	ORF IS231C [Bacillus thuringiensis]
713 Bt1Gc1013	Bt1G945	1794-1349	g2497392	754	753	1.20E-74	. 98	59	INSERTION SEQUENCE IS232 PUTATIVE ATP-
714 Bt1Gc1014	Bt1G946	1832-1	g2635858	278.	358	7.60E-32	38	26	(Z99121) yvgS [Bacillus subtilis]
714 Bt1Gc1014	Bt1G947	1770-1535	g732400	76	137	6.30E-09	34	29	HYPOTHETICAL 31.7 KD PROTEIN IN KINC-ADEC INTERGENIC REGION (ORF3) [Bacillus subtilis]
714 Bt1Gc1014	Bt1G948	2462-2141	g2619051	236	272	7.60E-23	52		(AF027868) RecQ homolog [Bacillus subtilis]
715 Bt1Gc1015	Bt1G949	135-548	g3676415	89	131	1.00E-08	22	99	(AF051917) unknown [Staphylococcus aureus]
716 Bt1Gc1017	Bt1G950	167-2407	g3130015	142	227	4.10E-18	26	49	(AL023517) putative iron-sulfur binding oxidoreductase [Streptomyces coelicolor]
717 Bt1Gc1016	Bt1G951	1-1596	g1171759	501	774	7.30E-77	35	93	BACILLOLYSIN PRECURSOR (NEUTRAL PROTEASE)
717 Bt1Gc1016	Bt1G952	1598-1864	£2696017	166	204	3.10E-15	4	= .	[Bacillus brevis] (AB001874) chitinase B [Clostridium paraputrificum]
718 Bt1Gc1018	Bt1G953	23-711	g4584144	949	717	8.00E-71	76	100	(AJ010140) DNA alkylation repair enzyme [Bacillus
719 Bt1Gc1019	Bt1G954	1-173	g2340003	156	130	1.30E-08	58	62	(Z98682) YlbG protein [Bacillus subtilis]
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(Z99110) similar to hypothetical proteins [Bacillus subtilis]	66		2.00E-12	166	110	g2633558	2153-647	Bt1G1014	751 Bt1Gc1057	_
INTERGENIC REGION [Bacillus subtilis] PROTEIN DLTD PRECURSOR [Bacillus subtilis]	29	53	1.10E-25	291	286	g729343	348-1	Bt1G1013	751 Bt1Gc1057	
thuringiensis [Bacillus thuringiensis] HYPOTHETICAL 80.1 KD PROTEIN IN SODA-COMGA	71	- 52	5.10E-124	1219	1198	g1731017	2535-609	Bt1G1012	750 Bt1Gc1052	_
[Bacillus subtilis] beta-lactamase (EC 3.5.2.6) precursor - Bacillus	86	95	1.10E-130	1282	1278	g1084221	803-1	Bt1G1011	750 Bt1Gc1052	
synthesis gene cluster of Streptomyces hygroscopicus										
(D50453) homologue of hypothetical protein in a rapamycin	100	39	1.10E-66	678	834	g1805444	1919-492	Bt1G1010	749 Bt1Gc1054 Bt1G1010	
synthesis gene cluster of Streptomyces nygroscopicus [Bacillus subtilis]										
(L)	9	38	9.30E-43	4/1	459	g1803443	4/4-1	BI101009	/49 BUIGETU34	
(Z99112) similar to processing protease [Bacillus subtilis]			1.10E-39	423	403	g2634058	1639-1158	Bt1G1008		
[Bacillus subtilis]	.*									
(Z99112) similar to 3-oxoacyl- acyl-carrier protein reductase	100	52	6.10E-57	586	613	g2634059	1054-344	Bt1G1007	748 Bt1Gc1051	
[Bacilling embrilie]	į				ļ	9				
[Bacillus subtilis] (Z99112) similar to hypothetical proteins from B. subtilis	99	54	1.30E-22	262	254	e2634060	265-11	Bt1G1006	748 Bt1Gc1051	
(D50453) homologue of unidentified protein of E. coli	36	52	3.60E-27	305	293	g1805397	1239-1560	Bt1G1005	747 Bt1Gc1050	
hypothetical protein P47K of P. chlororaphis [Bacillus subtilis]										
(DS0453) homologues to nitrile hydratase region 3'-	80	2	2.20E-100	996	1061	g1805408	950-1	Bt1G1004	747 Bt1Gc1050 Bt1G1004	
(SPORE GERMINATION PROTEIN C1) [Bacillus subtilis]									 	
SYNTHASE COMPONENT I (HEPPP SYNTHASE)									٠	
subtilis] PROBABLE HEPTAPRENYL DIPHOSPHATE	98	42	3.00E-32	353	370	g399774	778-1562	Bt1Ġ1003	746 Bt1Gc1053	
GTP CYCLOHYDROLASE I (GTP-CH-I) [Bacillus	99	79	5.50E-79	794	790	g127481	25-594	Bt1G1002	746 Bt1Gc1053	
(Z98682) YlbE protein [Bacillus subtilis]	99	4	1.80E-16	204	204	g2340001	1690-1454	Bt1G1001	745 Bt1Gc1049	
(Y14084) hypothetical protein [Bacillus subtilis]	99	38	1.70E-29	327	321	g2226254	1368-790	Bt1G1000	745 Bt1Gc1049	
(Z98682) YlbF protein [Bacillus subtilis]	99		6.50E-37	397	462	g2340002	678-235	Bt1G999	745 Bt1Gc1049	
thuringiensis] (Z98682) YlbG protein [Bacillus subtilis]	37	68	7.80E-09	132	132	g2340003	102-1	Bt1G998	745 Bt1Gc1049	
IMMUNE INHIBITOR A PRECURSOR [Bacillus	74	74	2.60E-200	1939	2075	g124464	1533-1	Bt1G997	744 Bt1Gc1048	
ALANYL-HISTIDINE DIPEPTIDASE) (CARNOSINASE) (PEPTIDASE D) [Haemophilus influenzae Rd]					•					
	:				Score					7
NCBI gi description	Cvrg	% % Ident Cvrg	BlastP- Prob	BlastP Score	nap	NCBI gi	Position	Gene Id	D Contig Id	
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SEQ ID Contig Id Gene Id NO 751 Bt1Gc1057 Bt1G1015 752 Bt1Gc1058 Bt1G1016 752 Bt1Gc1058 Bt1G1017 753 Bt1Gc1059 Bt1G1018 754 Bt1Gc1061 Bt1G1019 754 Bt1Gc1061 Bt1G1020 755 Bt1Gc1062 Bt1G1021 756 Bt1Gc1062 Bt1G1022	Position 2633-1275 198-1 1034-204 164-2198 78-1394 1481-1807 1273-750 1499-2618	NCBI gi g2632017 g3282095 g1709566 g1731017 g3023410 g2851549 g773349 g1788760		BlastP Score 830 227 992 1725 863 863 394 395	BlastP- Prob 8.50E-83 6.70E-19 5.80E-100 1.20E-177 2.70E-86 1.30E-36 1.10E-36 2.00E-08	% % 1dent Cvrg 38 82 59 23 71 100 53 93 50 100 56 53 49 32	% % % % % % % % % % % % % % % % % % %	(AJ002571) DppE [Bacillus subtilis] (AJ002571) DppE [Bacillus subtilis] (AJ007446) hypothetical protein [Thermotoga neapolitana] 3-METHYL-2-OXOBUTANOATE HYDROXYMETHYLTRANSFERASE (KETOPANTOATE HYDROXYMETHYLTRANSFERASE) [Bacillus subtilis] HYPOTHETICAL 80.1 KD PROTEIN IN SODA-COMGA INTERGENIC REGION [Bacillus subtilis] BRANCHED-CHAIN AMINO ACID TRANSPORT SYSTEM CARRIER PROTEIN (BRANCHED-CHAIN AMINO ACID UPTAKE CARRIER) [Bacillus subtilis] CSAA PROTEIN [Bacillus subtilis] (U20445) BirA protein [Bacillus subtilis] (AE000329) orf, hypothetical protein [Escherichia coli]
Bt1Gc1059 Bt1Gc1061	164-2198 78-1394	g1731017 g3023410	1593	1725 863	1.20E-177 2.70E-86	50	93 100	HYPOTHETICAL 80.1 KD PROTEIN IN SODA-COM INTERGENIC REGION [Bacillus subtilis] BRANCHED-CHAIN AMINO ACID TRANSPORT SYSTEM CARRIER PROTEIN (BRANCHED-CHAIN SYSTEM CARRIER PROTEIN (BRANCHED-CHAIN SYSTEM) (BRANCHED-CHAIN SYSTE
Bt1Gc1061	1481-1807	g2851549	382	394	1.30E-36	71	99	CSAA PROTEIN [Bacillus subtilis]
Bt1Gc1063 Bt1Gc1062	1273-750 1499-2618	g773349 g1788760	477 159	395 131	1.10E-36 2.00E-08	56 49	53 32	(U20445) BirA protein [Bacillus subtilis] (AE000329) orf. hypothetical protein [Escherichia coli
Bt1Gc1064	1021-221	g1881234	447	- 414	1.00E-38	35	100	(AB001488) FUNCTION UNKNOWN. [Bacillus subtilis]
757 Bt1Gc1064 Bt1G1024	1446-1653	g1722938	129	137	2.30E-09	39	51	HYPOTHETICAL 14.3 KD PROTEIN CY427.14 [Mycobacterium tuberculosis]
758 Bt1Gc1065 Bt1G1025	728-186	g2634575	133	191	4.40E-15	26	99	(Z99115) similar to phage-related protein [Bacteriophage SPBc2]
Bt1Gc1066	1-362	g3913081	469	472	7.30E-45	71	82	ARGININE REPRESSOR [Bacillus stearothermophilus]
	630-1962	g132256	1200	1187	1.30E-120	56	77	DNA REPAIR PROTEIN RECN (RECOMBINATION PROTEIN N) [Bacillus subtilis]
760 Bt1Gc1070 Bt1G1028 761 Bt1Gc1069 Bt1G1029	453-950 1699-1	g4584100 g1708643	806 439	811 453	8.70E-81 9.10E-43	91 33	99 52	(AJ010132) hypothetical protein [Bacillus cereus] (U51115) YebA [Bacillus subtilis]
761 Bt1Gc1069 Bt1G1030	1821-1205	g2829805	330	371	3.70E-34	39	75	HYPOTHETICAL 30.5 KD PROTEIN IN GABP-GUAA INTERGENIC REGION [Bacillus subtilis]
	1-1083	g2626818	874	675 200	2.30E-66	4	e 93	(D83967) YfkF [Bacillus subtilis]
763 Bt1Gc1074 Bt1G1033	1-704	g1001758	60	122	1.00E-07	ដូ ខ	37	(D64004) hypothetical protein [Synechocystis sp.]
764 Bt1Gc1072 Bt1G1034	863-1135	g2622856	228	243	1.40E-20	51	99	(AE000929) unknown [Methanobacterium thermoautotrophicum]
764 Bt1Gc1072 Bt1G1035	1466-2299	g1783243	1031	753	1.20E-74	69	100	(D83026) homologous to jojC gene product (B. subtilis; prf:2111327a); hypothetical [Bacillus subtilis]
765 Bt1Gc1073 Bt1G1036	1001-1	g135566	61	145	3.30E-09	29	21	TRYPSIN-RESISTANT SURFACE T6 PROTEIN PRECURSOR [Streptococcus pyogenes]

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Gene Id Position NCBI ginal BlastP Score BlastP Prob % % core BtIG1059 619-1738 g2634950 1140 1103 1.00E-111 59 100 BtIG1061 1932-688 g729608 1744 1752 1.70E-180 80 100 BtIG1062 740-1648 g1724002 762 732 2.10E-72 50 100 BtIG1063 1635-2579 g1724003 372 414 1.00E-38 30 100 BtIG1064 201-794 g3308998 473 516 1.60E-49 51 99 BtIG1065 1-1484 g1881239 140 228 4.20E-18 33 23 BtIG1066 1-1484 g1881239 140 228 4.20E-18 33 23 BtIG1067 1390-752 g586019 51 381 3.20E-35 48 100 BtIG1071 757-198 g2280497 100 124 2.50E-63 81 99
Gene Id Position NCBI ginary BlastP Score BlastP Prob % % core % % ldent Cvrg Score Blid Indent Cvrg Score Prob Ident Cvrg Ident Cvrg Score Brid Ident Cvrg Score Prob Ident Cvrg Ident Cvrg Score Prob Ident Cvrg Ident Cvrg Score Prob Ident Cvrg Ident Cvrg Ident Cvrg Ident Cvrg Score Prob Ident Cvrg Ident
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BlastP BlastP- % % Score Prob Ident Cvrg 1103 1.00E-111 59 100 638 1.90E-62 86 67 1752 1.70E-180 80 100 732 2.10E-72 50 100 414 1.00E-38 30 100 516 1.60E-49 51 99 431 2.50E-39 34 24 431 2.50E-39 34 24 228 4.20E-18 33 23 473 5.70E-45 37 100 299 1.60E-26 54 35 643 5.50E-63 81 99 124 2.50E-07 26 99 122 5.90E-11 37 99 123 7.30E-29 34 68 222 2.30E-18 40 90 185 1.90E-14 41 37 20
Ment Cyrg 59 100 86 67 80 100 50 100 51 99 51 99 54 35 81 99 54 35 68 40 99 54 37 74 100 54 39 54 39 54 39 55 40 56 49 57 40 58 40 59 54 59 54 50 54 51 99 51 99 52 64
Ment Cyrg 59 100 86 67 80 100 50 100 51 99 51 99 54 35 81 99 54 35 68 40 99 54 37 74 100 54 39 54 39 54 39 55 40 56 49 57 40 58 40 59 54 59 54 50 54 51 99 51 99 52 64
CVrg 100 100 100 100 100 337 37 100 39 99 99 64
NCBI gi description Z99116) similar to hypothetical proteins [Bacillus subtilizaciil phosphoribory phoribory proteins subtilis] AB015998) CBP21 precursor [Serratia marcescens] EXOGLUCANASE B PRECURSOR EXOCELLOBIOHYDROLASE B) (1,4-BETA-EXOCELLOBIOHYDROLASE B) (1,4-BETA-EXOCELLOBIOHYDROLASE B) (CBP120) [Cellulomonasimi] AB001488) FUNCTION UNKNOWN. [Bacillus subtilistypothetical] phoribory p
o hypothetical proteins [Bacillus subtitionIBOSYLTRANSFERASE (UMPRYLASE) (UPRTASE) [] XYMETHYLTRANSFERASE (SERILHMT) [Bacillus subtilis] ABC TRANSPORTER ATP-BINDIN LPD-CSPB INTERGENIC REGION LPD-CSPB INTERGENIC REGION J4.9 KD PROTEIN IN GLPD-CSPB [GION [Bacillus subtilis] 21 precursor [Serratia marcescens] E B PRECURSOR HYDROLASE B) (CBP120) [Cellulomonas OLASE B) (CBP120) [Cellulomonas CTION UNKNOWN. [Bacillus subtilis] PROTEIN H11664 [] PROTEIN H1664 [] PROTEIN M [Bacillus subtilis] ILATION PROTEIN M [Bacillus subtilis] OTEIN X [Bacillus subtilis] Insporter [Synechocystis sp.] family transcription regulator [Bacillus sp. SORY PROTEIN URED [Bacillus subtilis] SORY PROTEIN UREF [Bacillus sp. SORY PROTEIN UREF [Bacillus sp. SORY PROTEIN UREE [Bacillus sp.
teins [Bacillus subti NSFERASE (UMP ASE) [] NNSFERASE (SERII subtilis] RTER ATP-BINDIN RGENIC REGION [IN IN GLPD-CSPB subtilis] ratia marcescens] R (1,4-BETA- 120) [Cellulomonas WN. [Bacillus subtilis] 64 [] Sase [Escherichia collon protein HOMOLO PROTEIN HOMOLO PROTEIN HOMOLO PROTEIN Bacillus subtilis [Subtilis] lus subtilis] locystis sp.] on regulator [Bacillus subtilis] URED [Bacillus subtilis] UREG [Bacillus sp.] UREE [Bacillus sp.] UREE [Bacillus sp.] UREE [Bacillus sp.] UREE [Bacillus sp.] UREE [Bacillus sp.] UREE [Bacillus sp.]
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Table 1

(AB000617) YeeG [Bacillus subtilis]	%	41	4.4UE-86	861	847	g/415/44	220-1819	801 BEIGCIII3 BEIGII06	~
(AB000617) YeeF [Bacillus subtilis]	56		4.30E-24	276	431	g2415743	1-434	BtlGcl113	. ~-
(Z99117) similar to hypothetical proteins [Bacillus subtilis]	70	_	1.40E-107	_	1063	g2635199	857-1738	Bt1Gc1112.	. ~
(Z99117) similar to hypothetical proteins [Bacillus subtilis]	99	68	1.20E-33	366	349	g2635198	563-230	800 Bt1Gc1112 Bt1G1103	~
(Y09450) transposase [Pseudomonas putida]	29	30	2.80E-37	411	358	g1694898	1192-2078	799 Bt1Gc1108 Bt1G1102	
nyponencar 25:58 procen - Elicrococcus raccaris prasilità	100		4.400-43		1.1	817777		01100	_
binDing PROTEIN [insertion sequence IS232]	3	27	2 20E-20	326	2/12	a70050	044_300	700 Bt1C+1108 Bt1C1101	
INSEKTION SEQUENCE IS232 PUTATIVE ATP-	œ	91	5.10E-05	101	101	g249/392	Jan-66	/99 BUGGIOS BUGIO	
(AF024/13) ParC [Bacillus subtilis]			6.50E-25		324	g2558947	1454-1695	Brigerios	
TOPOISOMERASE IV SUBUNIT B [Bacillus subtilis]	74		5.20E-202	_	2092	g3914289	1-1452	Bt1Gc1109	
(AF065404) pXO1-138 [Bacillus anthracis]	99	_	7.30E-29		321	g4894353	985-1275	BtlGcllll	
beta-lactamase (EC 3.5.2.6) III precursor - Bacillus cereus []	<u>&</u>	_	7.50E-107	_	1059	g67770	2084-2868	Bt1Gc1110	
INTERGENIC REGION PRECURSOR [Bacillus subtilis]									ı
HYPOTHETICAL 13.0 KD PROTEIN IN IDH-DEOR	51	45	7.80E-09	132	89	g1731301	881-699	796 Bt1Gc1110 Bt1G1095	. 1
(AF027868) RAS-related protein [Bacillus subtilis]	43	28	1.50E-09		95	g2619009	289-1	796 Bt1Gc1110 Bt1G1094	
(ORF1) [Baçillus subtilis]									
HYPOTHETICAL 20.1 KD PROTEIN IN HMP 5'REGION	99	58	2.40E-46	486	491	g1730265	530-1	795 Bt1Gc1101 Bt1G1093	
horikoshii]									
(AP000007) 424aa long hypothetical protein [Pyrococcus	100	49	1.80E-82	827	943	g3258354	2172-860	794 Bt1Gc1095 Bt1G1092	- 1
acetyltransferase [Bacillus subtilis]									
(Z99110) similar to ribosomal-protein-alanine N-	83	26	1.40E-11	158	116	g2633543	1-451	794 Bt1Gc1095 Bt1G1091	_ 1
subtilis]									
(Z99110) similar to transcription regulation [Bacillus	99	51	2.90E-41	438	435	g2633560	1274-1750	793 Bt1Gc1106 Bt1G1090	. 1
[Helicobacter pylori J99]									
(AE001539) HISTIDYL-TRNA SYNTHETASE	72	35	3.90E-39	418	382	g4155714	942-1	793 Bt1Gc1106 Bt1G1089	
(AJ222587) YkuJ protein [Bacillus subtilis]	99	62	5.70E-22	256	247	g2632231	1310-1077	792 Bt1Gc1105 Bt1G1088	_,
(AJ222587) YkuK protein [Bacillus subtilis]	99		1.70E-45		469	g2632232	608-93	792 Bt1Gc1105 Bt1G1087	_ 1
CELL WALL HYDROLASE CWLJ [Bacillus subtilis]	99	57	4.60E-43	455	434	g1175629	1740-1311	791 Bt1Gc1102 Bt1G1086	. 1
(AE001167) lysyl-tRNA synthetase [Borrelia burgdorferi]	79	33	1.00E-55	553	493	g2688588	1-1250	791 Bt1Gc1102 Bt1G1085	_ 1
recE protein - Bacillus subtilis []	75	_	1.80E-89	893	991	g72992	782-1	Bt1Gc1100	_,
(Z99112) penicillin-binding protein [Bacillus subtilis]	51		4.20E-26	295	211	g2634067	699-1	Bt1Gc1099	. 1
INTERGENIC REGION [Escherichia coli]									
HYPOTHETICAL 138.1 KD PROTEIN IN MOLR-BGLX	17	31	1.40E-17	228	192	g2507582	2536-745	788 Bt1Gc1098 Bt1G1082	_1
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Contig Id		Bt1Gc1130	Bt1Gc1132	Bt1Gc1132	Bt1Gc1133		Bt1Gc1128	Bt1Gc1131			Bt1Gc1129	Bt1Gc1135	Rt1Gc1136		Bt1Gc1136	Bt1Gc1140	Bt1Gc1143	Bt1Gc1143	Bt1Gc1143	Bt1Gc1143	Bt1Gc1138	Bt1Gc1138	Bt1Gc1142	Bt1Gc1134	Bt1Gc1134	Bt1Gc1134		
Gene Id	•	Bt1G1128	Bt1G1129	Bt1G1130	Bt1G1131		Bt1G1132	Bt1G1133	B+1C1134	7	Bt1G1135	Bt1G1136	Rr1G1137 .		Bt1G1138	Bt1G1139	Bt1G1140	Bt1G1141	Bt1G1142	Bt1G1143	Bt1G1144	Bt1G1145	Bt1G1146	Bt1G1147	Bt1G1148	Bt1G1149		
Position		1026-1942	1159-938	1435-1	1-1026		1-855	147-1600	050 1063	607-1765	324-2325	1859-1	306-1		1746-1922	1018-1	1-273	298-795	812-997	1028-1139	775-1	1103-1609	1017-1	1-369	442-1047	1709-1872		
NCBI gi		g1076106	g2649348	g2634950	g1708267		g2465008	g2493592	2012212	80710210	g225714	g1213021	03257514	(g2633135	g3724051	g1172978	g1173265	g4512422	g4512423	g3688825	g1944618	g2851670	g2326729	ø1770039	g2621999		2/07/2
aat_		369	71	%	842		. 344	647	160	100	1796	578	146		132	. 813	364	748	221	158	638	285	952	155	126	128	î i	
BlastP Score		379	143	130	891		471	572) }		1388	477	143		135	773	364	748	226	96	677	327	850	176	169	145) } .	٥
BlastP- Prob		6.40E-35	5.30E-10	7.50E-08	2.90E-89		9.30E-45	1.90E-55	2 505 20	F.50E-20	7.40E-167	2.20E-45	4 30E-00		3.80E-09	9.30E-77	2.00E-33	4.10E-74	8.60E-19	5.10E-05	1.40E-66	1.70E-29	6.40E-85	1.70E-13	9 40E-13	1.20E-09	. Ž-van	× × ×
% % Ident Cvrg		32	38	သ	53		42	40	ş	2	54	4	3		47	4 6	79	90	73	95	51	38	56	32	24	47		2
%	(.)	54		23			77	100	3	É	86	63	<u>ر</u> د		.98	91	75	99	98	25	8	61	78	54	99	17		Š
NCBI gi description	[Methanococcus jannaschii]	schC protein - Streptomyces halstedii []	(AE001019) mutator protein MutT (mutT) [Archaeoglobus	(Z99116) similar to hypothetical proteins [Bacillus subtilis]	FLAVOHEMOPROTEIN (HAEMOGLOBIN-LIKE	PROTEIN) (FLAVOHEMOGLOBIN) [Bacillus subtilis]	(AJ001445) ripening-induced protein [Fragaria vesca]	HYPOTHETICAL 43.7 KD PROTEIN IN NPRE-PYCA	INTERGENIC REGION [Bacillus subtilis]	OXYGENASE SUBUNIT [Alcaligenes sp.]	urease [Canavalia ensiformis]	(X84710) ORF492, surface antigen gene [Methanosarcina	mazei] (A D000005) 403aa long hynothetical macrolide-effluy	determinant [Pyrococcus horikoshii]	(Z99108) yfjU [Bacillus subtilis]	(AJ000758) hypothetical protein [Bacillus megaterium]	50S RIBOSOMAL PROTEIN L18 [Bacillus subtilis]	30S RIBOSOMAL PROTEIN S5 (BS5) [Bacillus subtilis]	(AB017508) rpmD homologue (identity of 82% to B. subtilis) [Bacillus halodurans]	(AB017508) rplO homologue (identity of 84% to B. subtilis) [Bacillus halodurans]	(AF084104) NatA [Bacillus firmus]	(Y12602) acid phosphatase [Streptococcus equisimilis]	HYPOTHETICAL 49.0 KD PROTEIN IN BLTD-TRKA INTERGENIC REGION [Bacillus subtilis]	(Z98268) hypothetical protein Rv1686c [Mycobacterium	tuberculosis] (775208) hypothetical protein [Bacillus subtilis]	(AE000866) photoreactivation-associated protein		(299111) Similar to Spore cortex-lytic enzyme bacillus

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SEQ ID Contig Id Gene Id NO	Position	NCBI gi	nap Score	BlastP Score	BlastP- Prob	% % Ident Cvrg	% %	NCBI gi description
÷							•	subtilis]
828 Bt1Gc1145 Bt1G1151	479-39	g2632544	317	285	4.80E-25	-	99	(Z99105) ycbO [Bacillus subtilis]
828 Bt1Gc1145 Bt1G1152	2115-1675	g2632544	214	160	8.40E-12	33	99	(Z99105) ycbO [Bacillus subtilis]
829 Bt1Gc1144 Bt1G1153	2448-1366	g132565	946	982	6.60E-99	52	9	RIBOFLAVIN-SPECIFIC DEAMINASE [Bacillus subtilis]
830 Bt1Gc1150 Bt1G1154	230-988	g2117240	334	377	8.50E-35	34	100	(Z95586) hypothetical protein Rv1597 [Mycobacterium
830 Bt1Gc1150 Bt1G1155	1696-1216	g730088	118	207	8.80E-17	34	99	MUTATOR MUTT PROTEIN (7,8-DIHYDRO-8-
				•				OXOGUANINE-TRIPHOSPHATASE) (8-OXO- DGTPASE) (DGTP PYROPHOSPHOHYDROLASE)
								[Streptococcus pneumoniae]
830 Bt1Gc1150 Bt1G1156	2299-1851	g585393	311	332	5.00E-30	43	8 2	SIGNAL PEPTIDASE I P (SPASE I) (LEADER PEPTIDASE I) [Bacillus subtilis]
831 Bt1Gc1154 Bt1G1157	383-1	g4154045	140	189	9.20E-15	34	43	(AL035159) putative CDP-diacylglycerolserine O-
831 BtlGc1154 BtlG1158	1805-1927	g2224768	107	133	6.10E-09	60	57	phosphaticylicansierase [wycoodcienum ieprae] (Z97025) ylaf [Bacillus subtilis]
832 Bt1Gc1153 Bt1G1159	944-1	g417314	1561	1558	6.10E-160	_	94	MBL PROTEIN [Bacillus cereus]
832 Bt1Gc1153 Bt1G1160	1476-1108	g4584136	629	629	1.70E-61	99	99	(AJ010138) stage III sporulation protein D [Bacillus cereus]
833 Bt1Gc1146 Bt1G1161	985-170	g4753870	491	514	2.60E-49	3 8	100	(AL049754) putative oxidoreductase [Streptomyces coelicolor]
833 Bt1Gc1146 Bt1G1162	1668-1036	g1653142	248	296 ·	3.30E-26	33	100	(D90911) hypothetical protein [Synechocystis sp.]
833 Bt1Gc1146 Bt1G1163	2652-1704	g3913969	504	564	1.30E-54	36	71	KYNURENINASE (L-KYNURENINE HYDROLASE)
834 Bt1Gc1152 Bt1G1164	990-1	g1881291	294	3 81	3.20E-35	29	95	[Natius not regions] (AB001488) PROBABLE INTEGRASE. [Bacillus subtilis]
Bt1Gc1152	993-1	g166159	321	335	2.40E-30		95	
834 Bt1Gc1152 Bt1G1166	1451-1011	g4126639	271	292	8.70E-26	42	99	_
835 Bt1Gc1147 Bt1G1167	480-31	g1881290	510	431	1.60E-40	59	99	_
835 Bt1Gc1147 Bt1G1168	1566-860	g2632778	817	706	1.20E-69		30	_
836 Bt1Gc1148 Bt1G1169	269-1	g3930535	72	152	2.10E-09	32	. 7	(AF065313) DNA polymerase III; family C DNA
836 Bt1Gc1148 Bt1G1170	699-502	g2493764	306	306	2.90E-27	85	98	polymerase [Thermotoga maritima] COLD SHOCK-LIKE PROTEIN CSPD [Bacillus cereus]
836 Bt1Gc1148 Bt1G1171	1106-1816	g2649993	202	208	6.90E-17	35	100	(AE001061) conserved hypothetical protein [Archaeoglobus
836 Bt1Gc1148 Bt1G1172	1991-2222	g225559	360	376	1.10E-34	96	16	fulgidus] ORF IS231C [Bacillus thuringiensis]
Bt1Gc1151	1-941	g2632613	578	623	7.30E-61		92	

[Synechocystis sp.]					4, 5,				-	
PROBABLE ASPARAGINYL-TRNA SYNTHETASE (ASPARAGINE-TRNA I IGASE) (ASPARAGINE-TRNA I IGASE)	39 37		8.40E-31	341	244	g1711644	1192-1	Bt1G1193	848 Bt1Gc1171	
	,									
	58 100	_	8.20E-69	698	738	g466185	2223-1473	Bt1G1192	847 Bt1Gc1168	
INTERGENIC REGION (ORFX8) [Bacillus subtilis]						(
HYPOTHETICAL 22.0 KD PROTEIN IN RIBT-DACB	50 99	•	5.60E-47		451	g466186	1324-744	Bt1G1191	847 Bt1Gc1168	
(AB013369) ComEC [Bacillus halodurans]	27 73		8.80E-17	207	135	g4514331	586-1	Bt1G1190	847 Bt1Gc1168	
synthetase [Pyrococcus horikoshii]										
	29 26		1.30E-18	237	229	g3257481	1-853	Bt1G1189	846 Bt1Gc1165	
trachomatis]									•	
(AE001277) Isoleucyl-tRNA Synthetase: [Chlamydia	30 28		2.50E-23		314	g3328409	1-856	Bt1G1188	846 Bt1Gc1165	
(Z99120) similar to acyloate catabolism [Bacillus subtilis]	41 63	• -	2.00E-12	166	1:56	g2635785	1-251	Bt1G1187	845 Bt1Gc1164	
(AF015	48 100		3.50E-70		812	g2612898	1160-189	Bt1G1186	844 Bt1Gc1162	
_	49 100		9.20E-70		763	g2633815	1132-221	Bt1G1185	843 Bt1Gc1159	
(dihydrolipoamide acetyltransferase) [Bacillus subtilis]									-	
(Z99108) acetoin dehydrogenase E2 component	55 48		6.50E-53	548	54:4	g2633132	2983-2411	Bt1G1184	842 Bt1Gc1161	
subtilis]									•	
(AF006075) dihydrolipoamide dehydrogenase [Bacillus	57 100		6.70E-138	1350	1402	g2245640	2389-1016	Bt1G1183	842 Bt1Gc1161	
[Bacillus halodurans]				-						
(AB011836) similar to B.subtilis ywgB gene(27%-identity)			1.90E-14		. 149	g4512355	160-1	Bt1G1182	842 Bt1Gc1161	
(Y14080) hypothetical protein [Bacillus subtilis]	46 85		4.00E-110	1066	. 1160	g2226166	1-1488	Bt1G1181	841 Bt1Gc1163	
subtilis]	: .*					•				
(Z97025) product similar to E. coli PhoH protein [Bacillus	71 16	_			231	g2224770	1995-1778	Bt1G1180	840 Bt1Gc1160	
	37 99		3.30E-26		265	g2224771	1249-1740	Bt1G1179	840 Bt1Gc1160	
_	73 99		3.80E-32		352	g2224773	1090-812	Bt1G1178	840 Bt1Gc1160	
	<u>.</u>								•	
HYPOTHETICAL 43.7 KD PROTEIN IN NPRE-PYCA	34 58		9.40E-22	256	302	g2493592	680-1	Bt1G1177	840 Bt1Gc1160 Bt1G1177	
REDUCTASE (PEPTIDE MET(O) REDUCTASE)							•			
PUTATIVE PEPTIDE METHIONINE SULFOXIDE	65 56		1.20E-30	338	330	g1709688	1-298	Bt1G1176	839 Bt1Gc1157	
[Bacillus subtilis]										
(Z99111) similar to hypothetical proteins from B. subtilis	41 100	_	7.50E-43	453	607	g2633742	670-1715	Bt1G1175	838 Bt1Gc1156	
•						. (•	
GENERAL STRESS PROTEIN 20U (GSP20U) (DPS	59 97		5.20E-44	42	449	g3183517	420-1	Bt1G1174	838 Bt1Gc1156	
NCBI gi description	\circ		Prob	Score	nap . Score	NCR1 81	Position	Gene Id	NO Contigua	
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	Bt1Gc1185	Bt1Gc1185	Bt1Gc1184	Bt1Gc1180	Bt1Gc1183	Bt1Gc1183	71.001.10	Rt1Gc1183	Bt1Gc1182	Bt1Gc1177	Bt1Gc1177		Bt1Gc1176	Bt1 Gc11/6		Bt1Gc1178	Bt1Gc1178	Bt1Gc1175	•	Bt1Gc1175	Bt1Gc1169	Bt1Gc1169	Bt1Gc1173			R+1G-1173	Bt1Gc1172	849 Bt1Gc1172		Bt1Gc1171	Contig Id
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-	Bt1G1216	Bt1G1215	Bt1G1214	Bt1G1213	Bt1G1212	Bt1G1211	0	R+1G1210	Bt1G1209	Bt1G1208	Bt1G1207		Bt1G1206	Bt101205		Bt1G1204	Bt1G1203	Bt1G1202		Bt1G1201	Bt1G1200	Bt1G1199	Bt1G1198			R+1G1197	Bt1G1196	Bt1G1195		Bt1G1194	Gene Id
	1568-674	373-1	2223-695	1-641	2404-2210	1617-1993		40-1407	815-1545	1724-3056	1634-417		1276-884	85/-I		984-2096	1-418	2147-609	•	351-34	500-1501	198-1	2540-1293			1856-1	439-1116	Jan-88		1570-881	Position
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	g121839	g4584100	g3258180	g2635801	g2337807	g2635768	540000	3625346	e3845215	g2226188	g2226187	,	g141088	g2633600		g2649155	g1405459	g141186		g1881265	g2443235	g448837	g3915864	•	9,000	9730065	g1495280	g2735510	٠	g3123292	NCBI gi
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	1279	211	537	855	70	358	;	1370	4	678	812	·.·	185	484	5	155	121	72	٠.	376	1052	203	407	,	•		568	97	٠	866	
	1268	228	702	872	118	306	1200	1330	74	759	769		194	48/		250	131	123		380	1017	211	428		i	7773	599	106		605	BlastP Score
	3.30E-129	5.30	3.10	3.00	9.90	2.90	0.00	3 50E-135		2.80	2.50		2.10	1.90		2.40	1.00	7.00		4.10	1.30E-102	3.30	3.40		;	3 SOE_337	2 50	4.50		5.90	BlastP- Prob
	3-129	5.30E-19	3.10E-69	3.00E-87	9.90E-07	2.90E-27	1 1 2	13.5	0.72	2.80E-75	2.50E-76		2.10E-15	1.90년-46	;	2.40E-21	1.00E-08	7.00E-08		4.10E-35	-102	3.30E-17	3.40E-40		į	727	2 50E-58	4.50E-06		5.90E-59	<u> </u>
	80	39	35	78	34	55	۶	<u>د</u> د	26	37	40		34	S S	;	32	27	42		58	58	59	48		č	7,	<u> </u>	67		55	% Ident
	2	74	100	46	22	99	į	3	2	76	100		99	y		70	72	37	•	99	100	32	63		(80	<u>=</u>	53		28	% Cvrg
	64 ENDOGLUCANASE PRECURSOR (ENDO-1,4-BETA-GLUCANASE) (CELLULASE) (ENDO-K) [Bacillus sp.]	dehydrogenase [Pyrococcus horikoshii] 74 (AJ010132) hypothetical protein [Bacillus cereus]	_		_	(Z9912)		(70011				3'REGION (ORFD) [Clostridium perfringens]	subtilisj 99 HYPOTHETICAL 14.9 KD PROTEIN IN NAGH		[Archae				subtilis]					(SWIVELASE) [Bacillus subtilis]			dehydrase homolog [Staphylococcus carnosus] 00 (Z71928) hynothetical protein [Bacillus subtilis]		(PHENYLALANINEIRNA LIGASE BETA CHAIN) (PHERS) [Bacillus subtilis]		rg NCBI gi description
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SEQ NO	861]		861 1	861]		862	863			863 1	864	865]	866 1	866]	867				808	869	870]	870]	871 1	872]	872]	873]	
Contig Id	Bt1Gc1186		3t1Gc	Bt1Gc1186		Bt1Gc1188	B+1G-1188	Rt1Gc1197		3t1Gc	Bt1Gc1189	Bt1Gc1190	Bt1Gc1193	Bt1Gc1193	Bt1Gc1191	5	B11/C1191	Bt1Gc1195	PELIOCITAS	i Gc	Bt1Gc1198	Bt1Gc1198	Bt1Gc1200	Bt1Gc1199	Bt1Gc1199	Bt1Gc1197	
ig Id	1186		Bt1Gc1186	1186		1188	·1188	1193		Bt1Gc1192	:1189	:1190	:1193	1193	1191	1101	1191	1195		Bt1Gc1196	:1198	:1198	:1200	1199	:1199	:1197	
Ge	BtlC		Bt10	Bt10		Bt10	₽ <u>+</u>	P :	}		Bt10	Bt10	Bt10	Bt10	Btl	D+1	Bild	Bti	9118	Btl	Bt10	BtlC	Bt10	Bt10	Bt10	Bt10	
Gene Id	Bt1G1217		Bt1G1218	Bt1G1219		Bt1G1220	B+1G1331	Rt1G1222	i	Bt1G1223	Bt1G1224	Bt1G1225	Bt1G1226	Bt1G1227	Bt1G1228	11330	BII G 1229	Bt1G1230	Bt1G1231	Bt1G1232	Bt1G1233	Bt1G1234	Bt1G1235	Bt1G1236	Bt1G1237	Bt1G1238	
Po:	-		367	108	-	70-	153	141	:	2109	÷	688	16	160	25	Š	. 223	} -	/80	452		100	584	<u>.</u>	864	-	
Position	1-360		367-1037	1084-1848		70-1335	526-1000	1413-185	,	2109-1564	1-151	688-2037	1689-1	1608-721	25-807	1315	323-1210	1-472	/80-1594	452-2266	1-682	1001-1291	584-1735	1-830	864-1812	1-1516	
Z	g25		g11	g11	(g28	a 1 4	2 2 2	ď	g29	£19	g13	g42	<u>82</u> 1	g11	2	2 G2	g26	g24	g58	<u>9</u> 10	g21	g17.	g45	g26	g28	
NCBI gi	g2500507		g1168648	g1168649		g2827439	0715	61800301		g2982980	g1945675	g1303913	g4234795	g2116764	g1170788	2007	g/30171	g2633171	g2495571	g586060	g1002992	g2160199	g1730958	g4589056	g2635903	g2815318	
aat_ nap Score	. 4 .		1120	1033		1774	ስ)))	t		=	1932	ယ္လ	پې	4.	<u>ي</u>	بر د	·μ		2374	2	_	1427	1132	1282	7	
	482			33 -			<u>.</u>	71	,	127	124		320	338	496	h h	32.0	324	95		208	111				718	
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BlastP- Prob	6.4		1.10	1.1		7.80	7	۸ ر د د	į	6.6	7.10	4.40]	2.5	1.0	7.10	7	4.	2.6	2.3	9.801	4.3	9.0	3.40]	8.60	2.10	4.8	
•	6.40E-46		1.10E-114	1.10E-96		7.80E-183) i	5 50E-24	ì	6.60E-12	7.10E-10	4.40E-189	2.50E-35	1.00E-38	7.10E-47	1	4./0E-34	2.60E-33	2.30E-18	9.80E-240	4.30E-24	9.00E-08	3.40E-143	8.60E-106	2.10E-127	4.80E-73	
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		Bac	MENA		CYT	(AF	Visco		pneu	(AE	(Z94	Ď	(AF	Œ.	9	(PLP3)	AIP-B	(Z99	influ	ME	TRNA (U347)	ŒI:	HYPO	(AF	phos (Z99	<u> </u>	COCIICO
	NAQI R R	SOLFOR SOBONII (KIESKI [Bacillus stearothermophilus]	VAQU	MENAQUINOL-CYTOCHROME C REDUCTASE	CYTOCHROME B/C SUBUNIT [Bacillus subtilis]	(AF043609) aluminum resistance protein [Arthrobacter	viscosus]	NEOCLATON I FNOTEIN OLINE [Bacillus cereus]	pneumoniae]	(AE000682) transc	(Z94043) hypothetical protein [Bacillus subtilis]	(D84432) YqhX [Bacillus subtilis]	(AF078135) unknown [Leptospira borgpetersenii]	(D86418) YfnF [Bacillus subtilis]			E E	108)	HYPOTHETICAL PROTEIN HI0105 [Haemophilus influenzae Rd]	OIH		(D13377) XpaF1 protein [Bacillus licheniformis]	E OT	(AF120090) 2,3-bisphosphoglycerate-independent	phosphoglycerate mutase [Bacillus megaterium] (Z99121) enolase [Bacillus subtilis]	(AL021529) putative export protein [Streptomyces	Loron
		stearc	QUINOL-CYTOCHROME C REDUCTASE		RON	9) alı	aCT.		ae]	(2) (1)	hypo	Yqh	5) un	YfnF	EM	Pasteurella haemolytical	INDING PROTEIN ABC [Escherichia coli]	simil	ETIC e Rd]	XX	LIGASE) (METRS) [Bacillus subtilis] 2) ORF375 [Dichelobacter nodosus]	Xpal	EIIC	0)2,	lycera enola	19) pt	_
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	Bt1G1259	Bt1G1258	Bt1G1257	Bt1G1256	Bt1G1255	Bt1G1254	Bt1G1253		Bt1G1252	Bt1G1251			Bt1G1250		Bt1G1249	Bt1G1248	Bt1G1247		Bt1G1246	Bt1G1245		Bt1G1244		Bt1G1243	Bt1G1242	B1101241		Bt1G1240		Bt1G1239	Gene Id	Š	
	1782-1086	959-39	3163-681	1852-1211	1-1008	1-2170	1342-218		963-2056	575-1			1279-1645		388-984	1-308	1656-1236		1355-1	65-1612		1057-1257		96-1016	2186-1862	C/ 41-42/		207-1		1993-2142	Position		
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	g1730193	g400209	g3821797	g551727	g1934838	g1171128	g1945051		g1708975	g133949			g585920	-	g586914	g71129	g2633808		g2633180	g4835822		g141471		g1894757	g1620923	goodyyy	355000	g1730937	Q	g1345692	MCBI 8		
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	1042	688	233	896	675	1416	409		860	846			593		545	156	379		135	1146		250		902	142	023	3	176		156	Score	BlastP	
	2.90E-105	9.4(9.7(8.60	2.30	6.80E-145	3.5(5.60	1.70			1.30		1.30	2.20	5.20		3.80	2.80E-116		2.40		2.00	1.80	٠.)د	1	1.70	!	2.20	Prob	BlastP-	1
	3-105	9.40E-68	9.70E-19	8.60E-90	2.30E-66	<u>1</u> -145	3.50E-38		5.60E-86	1.70E-84		•	1.30E-56		1.30E-52	2.20E-11	5.20E-35		3.80E-09	3-116		2.40E-21		2.00E-90	1.80E-09	10-30E-01) }	1.70E-13		2.20E-11		'P	•
	83	51	35	80	43	42	39		45	8 2			92		2	77	62		32	48		74		58	30	. 5	.	4	;	53	Ident Cvrg	%	
	68	100	35	70	74	42	100		93	96		•	10		100	85	35		67	100		36		<u>1</u> 00	39	4	2	35	. ;	23	Cvrg	%	
(GA)	UDP	MEN	(D10	(M5)	(Z93940	(024	(163	[Pset	MET	30S 1	BETA	(TRA	DNA	ITINI	НҮРО.	ribos	(Z99	Bac	mone (Z99	(AF1	PHO	TRA	subtilis	(Z92	(Z79)	(ALU3	Bac	НУРО	ָרָרָ בַּרְי	CHLO			
(GALACTOWALDENASE) (UDP-GALACTOSE 4-EPIMERASE) [Bacillus subtilis]	UDP-GLUCOSE 4-EPIMERASE	ŒRΑ	(D10594) chitinase D precursor [Bacillus circulans]	(M57689) sporulation protein [Bacillus subtilis]	xantnusj (Z93940) unknown [Bacillus subtilis]	(U24657) saframycin Mx1 synthetase B [Myxococcus	(U63928) L1 protein [Bacillus cereus]	[Pseudomonas putida]	LOIH	RIBO	A SUI	NSC	-DIR	RGE	HIO	ribosomal protein L7/L12 - Bacillus subtilis []	111) s	[Bacillus subtilis]	monocytogenes] (Z99108) similai	(AF102174) glycine betaine transporter Betl [Listeria	SPHC	TRANŠALDOLASE-LIKE PROTEIN (20 KD	ES.	953) p	coencolor] (Z79580) putative ORF [Bacillus subtilis]	3131	[Bacillus subtilis]	HYPOTHETICAL 21.6	1.	OR A			
SE)	COS	NE-E	chitin	nods	unkno	safran	L1 pr	onas p	NE NE	MOS	SUBUNIT) [Bacillus subtilis]	RIPT	ECTE	NIC	ETIC/	protei	simila	ubtili	genes] simila	4) gly)PRO	DOL	,	peripl	outativ) nyi	ubtilis	ETIC.	m h	MPHI			
LDE Bacil	E 4-E	Š	ase D	lation	Œ.	nycin	otein	utida	GAM	AL PI	E E	ASE I	D R	EGI	1 L 22	n L7/	r to h	ٽ	r to h	cine t	TEIN	ASE-		asmic	⁄e OR	юше	<u>ب</u>	\L' 21		NIC	7	Z	
lus su	PIME	D PR	precu	prote	Bacill	MxI	[Baci	_	MA-I	OTE	a cill	3ETA	A PC	E N N	.5 KI	L12 -	poth		poth	etain	ORF	LIKE		subst	편 파	icai p	•	.6 KI	3	A JC	NCBI gi description) 	
E) (UI Ibtilis	RASI	OTE	LSOL	E E	us sul	synth	llus c		LYAS	N S	s sub	E C H	٨¥٦	23) (PRC	Bacil	etical	•	etical	e tran	<u>ე</u>	PRO		rate-b	acillu	готеш	•) PRC	. !	CETY	i desc		
DP-G]	(1)	N C	Baci	3acill ı	btilis]	etase	ereus) E	#(BS	tilis]		IERA	ORF2	TEN	llus su	prote		prote	sporte	S19)	TEIN		indin	s subt	200		TEI		LIR	indu		
ALA		TR	llus c	dus st		<u>Β</u> [Ν	_		MET	4) [B	·	RNA	SE B	3) 日	ZZ	ıbtilis	ins [I		ins fr	r Bet	[Baci	(20.1		g pro	ilis]	C / .Uo		N		ANSF	. 6	ì	
CTOS		Bacil	ircula	tilis]		yxoc	,		HION	acillu		POL	ETA (lacillu	PLL.		Bacill		om B.	LEi	llus sı	Ĝ		ein [lour of	5	LVA		ERA			
E 4-		MEMBRANE-BOUND PROTEIN LYTR [Bacillus subtilis]	ns]			occus			METHIONINE GAMMA-LYASE (L-METHIONINASE)	30S RIBOSOMAL PROTEIN S4 (BS4) [Bacillus subtilis]		(TRANSCRIPTASE BETA CHAIN) (RNA POLYMERASE	DNA-DIRECTED RNA POLYMERASE BETA CHAIN	INTERGENIC REGION (P23) (ORF23) [Bacillus subtilis]	THETICAL 22.5 KD PROTEIN IN RPLL-RPOB		(Z99111) similar to hypothetical proteins [Bacillus subtilis]		monocytogenes] (Z99108) similar to hypothetical proteins from B. subtilis	steria	PHOSPHOPROTEIN ORFU) (CSI9) [Bacillus subtilis]			(Z92953) periplasmic substrate-binding protein [Bacillus		1515) nypomencai protem scac7.08 (streptomyces	•	THETICAL 21.6 KD PROTEIN IN ILVA 3'REGION		RAMPHENICOL ACETYLTRANSFERASE (CAT)	•		
		btilis]			."		•	-	Ě	ilis]		WASE	Z	ilis]	w	•	tilis]		si		_			us		yces		NOI	- (E E			

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915 Bt1Gc1244 Bt1G1322	915 Bt1Gc1244 Bt1G1321	914 Bt1Gc1242 Bt1G1319 914 Bt1Gc1242 Bt1G1320	913 Bt1Gc1245 Bt1G1318	911 Bt1Gc1241 Bt1G1316 912 Bt1Gc1240 Bt1G1317	910 Bt1Gc1235 Bt1G1315	910 Bt1Gc1235 Bt1G1314	909 Bt1Gc1239 Bt1G1313	909 Bt1Gc1239 Bt1G1312	909 Bt1Gc1239 Bt1G1311	908 Bt1Gc1237 Bt1G1310	907 Bt1Gc1236 Bt1G1308 908 Bt1Gc1237 Bt1G1309	907 Bt1Gc1236 Bt1G1307	906 Bt1Gc1229 Bt1G1305 907 Bt1Gc1236 Bt1G1306	H	SEQ Contig Id Gene Id
2017-1	701-150	169-1004 1880-1315	2490-1765	1488-1 2188-140	3033-2173	3187-1	2141-2874	1643-723	626-1	1235-2173	3440-2581 571-1116	2219-1278	1-1778 1-752	1-1010	Position
g4680703	g729934	g2226222 g418449	g1894750	g2619051 g1072970	g1894770	g2633724	g4584149	g4584150	g4584151	g1731068	g769829 g2633543	g585208	g2182990 g2495393	g2769532	NCRI oi
304	446	351 126	744	1586 618	401	562	1234	1603	1020	1248	469 330	745	335 358	Score 226	
352	477	434 153	759	1489	440	624	1173	1498	738	1213	505 351	832	377 439	Score 280	BlastP
3.80E-32	2.20E-45	7.80E-41 4.70E-11	2.80E-75	1.20E-152 1.20E-65	1.80E-41	5.70E-61	3.80E-119	1.40E-153	4.80E-73	2.20E-123	2.30E-48 4.90E-32	5.20E-83	8.50E-35 2.30E-41	Prob 1.60E-24	BlastP-
34	49	34 24	60	6 4	36	48	98	100	98	78	37 38	52	32 39	Ident (%
88	99	100 99	100	84 78	100	35	78	100	92	%	99	100	53 85	Cyrg 61	%
(AF132966) CGI-32 protein [Homo sapiens]	[Bacillus subtilis] SIGNAL PEPTIDASE I (SPASE I) (LEADER PEPTIDASE D. GOGILLO GOLDANO	(Y14082) hypothetical protein [Bacillus subtilis] HYPOTHETICAL TRANSCRIPTIONAL REGULATOR IN HEMY-GLTT INTERGENIC REGION (ORFA)	autotrophicus] (Z92952) product similar to E.coli YjaF protein [Bacillus	proteins and to E. coli YidA protein [Bacillus subtilis] (AF027868) RecQ homolog [Bacillus subtilis] dhlR protein - Xanthobacter autotrophicus [Xanthobacter	[Bacıllus subtilis] (Z92954) product similar to Bacillus subtilis YxeH and YcsE	(Z99111) similar to two-component sensor histidine kinase	(AJ010111) cytochrome aa3 controlling protein [Bacillus	cereus] (AJ010111) cytochrome caa3 oxidase assembly factor	(AJ010111) cytochrome caa3 oxidase subunit II [Bacillus	acetyltransferase [Bacillus subtilis] HYPOTHETICAL 39.7 KD PROTEIN IN GLNQ-ANSR	(M80674) hydroxylase [Streptomyces glaucescens] (Z99110) similar to ribosomal-protein-alanine N-	sp.] GLYCEROPHOSPHORYL DIESTER PHOSPHODIESTERASE (GLYCEROPHOSPHODIESTER PHOSPHODIESTERASE) [Bacillus subtilis]	Japonicum] (U81485) histidine kinase [Lactococcus lactis cremoris] TRANSCRIPTIONAL REGULATOR SOXR [Arthrobacter	(AJ223073) phosphate regulatory protein [Bradyrhizobium	NORI ai Appariation

926	925	924]	924		924	923]	F)	923 1	922	922	922	921]	921	920		920]	919 1			918 1	, 710			917 1		917 1	916 1	916 1	915 1	NO BEQ
Bt1Gc1259	Bt1Gc1258	Bt1Gc1257	Bt1Gc1257		Bt1Gc1257	Bt1Gc1256		Bt1Gc1256	Bt1Gc1253	Bt1Gc1253	Bt1Gc1253	Bt1Gc1254	Bt1Gc1254	Bt1Gc1251		Bt1Gc1251	Bt1Gc1247		Bt1Gc1249	Bt1Gc1249	247130119	0+1C2-1240		Bt1Gc1248		Bt1Gc1248	Bt1Gc1246	Bt1Gc1246	Bt1Gc1244	Contig Id
Bt1G1345	Bt1G1344	Bt1G1343	Bt1G1342		Bt1G1341	Bt1G1340		Bt1G1339	Bt1G1338	Bt1G1337	Bt1G1336	Bt1G1335	Bt1G1334	Bt1G1333		Bt1G1332	Bt1G1331		Bt1G1330	Bt1G1329	07010130			Bt1G1327		Bt1G1326	Bt1G1325	Bt1G1324	Bt1G1323	Gene Id
1871-671	2021-1308	1420-1031	1025-426		400-1	2827-3336		507-2318	846-2229	351-710	1-223	2440-1284	1-1020	2062-1790	•	1799-1	172-1981		2542-2008	1947-1105	1-6201	1006 1		2096-2288		698-411	2416-2724	19-966	2302-1625	Position
g1731041	g1176948	g322197	g1770060	•	g2501608	g1750115		g1296829	g2500757	g620085	g2738159	g3913040	g2649875	g4126631		g2633600	g4151933	(g1805397	g1805396	82470033	~7406675		g4096799	(g113009	g2293270	g400209	g1881346	NCBI gi
119	536	379	547		362	769		1041	1130	441	153	771	54	129	i	482	653		329	244	202		•	2		265	243	516	527	nap Score
· 181	549	406	499			772		1149	1124	454	171	850	126	128	į	485	795			275	#/1	471		112		253	244	٠.	528	BlastP Score
5.00E-14	5.10E-53	7.20E-38	1.00E-47		8.50E-35	1.20E-76		1.30E-116	5.90E-114	5.90E-43	5.80E-13	6.40E-85	3.40E-08	2.10E-08		3.10E-46	4.30E-79		2.30E-34	5.50E-24	/.00E-4+	7 60E 44		1.00E-06		1.20E-21	1.10E-20	2.20E-52	8.50E-51	BlastP- Prob
1 34	43	5. 62	57		51	82		42	59	74	46	43	28	30			34			27	4			31	_	49	4. 48	42	45	% Ident
99	100	53	99		78	99		100	67	. 99	. 28	99	72	67		54	100	-	క్ష	100	. 0	2	200	.36		99	4 3	100	100	% Cvrg
INTERGENIC REGION [Bacillus subtilis] HYPOTHETICAL 14.6 KD PROTEIN IN GCVT-			_				lactis]		CHEMOTAXIS PROTEIN CHEA [Listeria		(U91841) MotB homolog [Bacillus firmus]	[Archaeoglobus fulgidus] PUTATIVE ALANINE RACEMASE [Bacillus subtilis]	? (AE001054) ribosomal protein S18 alanine acetyltransferase			_	_		•	sp. NGK234]) (D50453) ycgQ [Bacillus subtilis]	niro	carnosu	conceptual translation supplied by author. [Staphylococcus	(U4015		_		MEMB	(AB001488) FUNCTION UNKNOWN. [Bacillus subtilis]	NCBI gi description

70			Casa		NCBI C.	aat_]	BlastP	BlastP-	%	%	
	ON	Q			. 0	Score	Score	Prob	Ident Cvrg	91V	0
					,						SPOIIIAA INTERGENIC REGION [Bacillus subtilis]
	926 Bt	Bt1Gc1259	Bt1G1346	2085-1297	g1707680	365	497	1.60E-47	3 8	100	(Y08256) orf c06001 [Sulfolobus solfataricus]
	926 Bt	Bt1Gc1259	Bt1G1347	3345-2111	g1652676	971	1042	2.90E-105	47		(D90907) hypothetical protein [Synechocystis sp.]
. •			Bt1G1348	1-2428	g128494	72	180	1.20E-12	22		NODULATION PROTEIN V [Bradyrhizobium japonicum]
		Bt1Gc1261	Bt1G1349	1-2428	g2633696	747	746	6.70E-74	36		(Z99110) similar to hypothetical proteins [Bacillus subtilis
			Bt1G1350	534-1160	g2226262	316	361	4.20E-33	39	_	(Y14084) competence transcription factor [Bacillus subtilis]
	928 Bt		Bt1G1351	1480-2101	g2226177	536	559	4.40E-54	49		(Y14081) hypothetical protein [Bacillus subtilis]
	929 Bt	Bt1Gc1260	Bt1G1352	340-2256	g135176	2316	2231	2.90E-231	66		THREONYL-TRNA SYNTHETASE 2 (THREONINE-
					1						TRNA LIGASE) (THRRS) [Bacillus subtilis]
	929 Bt	Bt1Gc1260	Bt1G1353	2515-3146	g2293177	564	582	1.60E-56	51	83	(AF008220) transporter [Bacillus subtilis]
	930 Bt	Bt1Gc1263	Bt1G1354	1121-297	g2499116	335	371	3.70E-34	29	100	VANCOMYCIN B-TYPE RESISTANCE PROTEIN
			,			٠				_	VANW [Enterococcus faecalis]
-	930 Bt	Bt1Gc1263	Bt1G1355	2867-2104	g143324	998	998	1.30E-100	74	99,	(M37169) APase I [Bacillus licheniformis]
	931 Bt	Bt1Gc1264	Bt1G1356	1-905	g887872	203	355	1.80E-32	35	82	(U25453) phosphotidyl inositol-specific phospholipase C
					•						[Listeria monocytogenes]
٠	931 Bt	Bt1Gc1264	Bt1G1357	2171-1447	g116300	339	423	4.30E-39	40 ′	34 (CHITINASE A1 PRECURSOR [Bacillus circulans]
	931 Bt	Bt1Gc1264	Bt1G1358	2781-2179	g3308998	465	516	1.60E-49	51	99	(AB015998) CBP21 precursor [Serratia marcescens]
	932 Bt	Bt1Gc1262	Bt1G1359	1-399	g3183496	239	253	1.20E-21	35	65	HYPOTHETICAL 23.4 KD PROTEIN IN AAPA-SIGV
					٠.						INTERGENIC REGION [Bacillus subtilis]
	932 Bt	Bt1Gc1262	Bt1G1360	1936-470	g1170977	1464	1378	7.20E-141	57	100	PROBABLE METHYLMALONATE-SEMIALDEHYDE
						-					DEHYDROGENASE [Bacillus subtilis]
	932 Bt	Bt1Gc1262	Bt1G1361	2865-1972	g140739	407	477	2.20E-45	35	95	HYPOTHETICAL 31.0 KD PROTEIN IN RNPB-SOHA
											INTERGENIC REGION (ORF 2) [Escherichia coli]
	932 Bt	Bt1Gc1262	Bt1G1362	3925-2873	g2635778	641	560	3.50E-54	43	83	(Z99120) similar to butyryl-CoA dehydrogenase [Bacillus subtilis]
	933 Bt	Bt1Gc1266	Bt1G1363	1128-1520	g2633504	571	571	2.40E-55	<u>8</u> 1	99	(Z99110) similar to hypothetical proteins [Bacillus subtilis]
	933 Bt	Bt1Gc1266	Bt1G1364	2247-1591	g2633505	550	396	8.30E-37	49		(Z99110) similar to hypothetical proteins [Bacillus subtilis]
	934 Bt	Bt1Gc1267	Bt1G1366	750-1	g1170997	361	166	2.40E-12	33	93	HYPOTHETICAL 29.3 KD PROTEIN IN CCPA
											3'REGION (ORF1) [Bacillus megaterium]
٠	934 Bt	Bt1Gc1267	Bt1G1367	1780-1598	g4894352	159	12	3.20E-12	52	98	(AF065404) pXO1-137 [Bacillus anthracis]
	934 Bt	Bt1Gc1267	Bt1G1368	3057-3401	g2467222	8 2	113	6.40E-06	25	27	(X92946) macrolide efflux protein [Lactococcus lactis]
	935 Bt	Bt1Gc1269	Bt1G1369	1009-107	g1168646	1210	1144	4.50E-116	74	100	GTP-BINDING PROTEIN ERA HOMOLOG (BEX
	935 Rt		Br1G1370	1420-830	g1397290	248	300	9 40E-29	3		PROTEIN) [Bacillus subtilis]
		p(1001207	p(101370	1429-630	81377290	042	020	9.40E-29	1	7	(001949) Similar to cytome deaminase. [Caenornaodins elegans]

[Bacillus subtilis] GTP-BINDING PROTEIN TYPA/BIPA HOMOLOG [Bacillus subtilis]	22	5 77	1.50E-46	488	517	g3183185	1-398	Bt1Gc1277 Bt1G1389	942 Bt1Gc1
ALANYL-D-ALANINE CARBOXYPEPTIDASE) (DD-PEPTIDASE) (DD-CARBOXYPEPTIDASE) (PBP-5*)				•		4 .			
INTERGENIC REGION PRECURSOR [Bacillus subtilis] PENICILLIN-BINDING PROTEIN 5* PRECURSOR (D-	8 1	33	3.10E-37	400	317	g461914	1144-1	278 Bt1G1388	941, Bt1Gc1278
	100	30	2.50E-35	382	334	g1730928	1630-851	276 Bt1G1387	940 Bt1Gc1276
	100	•	1.10E-36	•	516	g1934806	737-63	276 Bt1G1386	940 Bt1Gc1276
	100	_	1.50E-257	2479	2621	g1339850	24-1874	273 Bt1G1385	939 Bt1Gc1273
subtilis] (Y13937) putative PacL protein [Bacillus subtilis]		60	1.80E-54	568	581	g2337795	2656-3247	268 Bt1G1384	938 Bt1Gc1268
(AJ000974) putative fibronectin-binding protein [Bacillus	100	62	1.70E-164	1601	1851	g2462963	2556-841	268 Bt1G1383	938 Bt1Gc1268
INTERGENIC REGION [Bacillus subtilis]						. (
PUTATIVE PEPTIDASE IN GCVT-SPOIIIAA	47	49	6.50E-37	397	378	g1731048	490-1	268 Bt1G1382	938 Bt1Gc1268
CHAI		£	7.5VE-12	107	131	8402404	2740-2720	7/2 DITO1301	237 BHOG1272
	1 9		2.70E-22	259	240	g141186	2564-2034		937 Bt1Gc1272
acetyltr					· :				
	99		2.60E-17	212	160	g2633543	2558-2034	272 Bt1G1379	937 Bt1Gc1272
_	52	53	2.60E-42	448	430	g2633171	462-1	272 Bt1G1378	937 Bt1Gc1272
INTERGENIC REGION [Bacillus subtilis]	-					. •			٠
	98	70	2.30E-16	203	203	g1730934	2487-2320	270 Bt1G1377	936 Bt1Gc1270
INTER					٠.,				
stearothermophilus] HYPOTHETICAL 17.9 KD PROTEIN IN DING-ASPB		သ အ	4.20E-17	210	188	g1730935	2315-1815	Bt1Gc1270 Bt1G1376	936 Bt1Gc1;
(TRANSAMINASE A) (ASPAT) [Bacillus			٠		14.5				•
	100		1.50E-154	1507	1522	g2492839	1787-612	270 Bt1G1375	936 Bt1Gc1270
DNA REPLICATION PROTEIN DNAD [Bacillus subtilis]	69	45	1.20E-33	366	394	g729356	482-1	270 Bt1G1374	936 Bt1Gc1270
INTERGENIC REGION [Bacilling subtilis]	. ,			;		0	1		
INTERGENIC REGION [Bacillus subtilis] HYPOTHETICAL 79.2 KD PROTEIN IN PHOH-DGKA	LA.	63 -	7.60E-08	134	117	g1730999	2443-2326	269 Bt1G1373	935 Bt1Gc1269
	99	2	9.80E-34	367	528	g1731000	2320-1850	269 Bt1G1372	935 Bt1Gc1269
	99	49	3.00E-16	202	276	g1708628	1847-1506	269 Bt1G1371	935 Bt1Gc1269
NCBI gi description	Cyrg	% Ident	BlastP- Prob	BlastP Score	nap Score	NCBI gi	Position	Id Gene Id	ID Contig Id
					•	-			CEO

					,	•			
[Cloning vector pr. W 13] 9 (U93876) hypothetical protein YrdC [Bacillus subtilis] 9 (AE000245) alcohol dehydrogenase [Escherichia coli]	32 99 66 19	1.30E-17 4.00E-18	215 221	179 220	g1934645 g1787753	686-1675 2294-2094	83 · Bt1G1409 85 · Bt1G1410	950 Bt1Gc1283 - 951 Bt1Gc1285	
	41 42	1.80E-34	374	374	g1277135	121-2539	83 Bt1G1408	950 Bt1Gc1283	
	48 ; 30	1.00E-45	480	446	g2635778	2642-3186	81 Bt1G1407	949 Bt1Gc1281	
	73 100	1.50E-110	1092 1	1139	g1731059	1587-2489	81 Bt1G1406	949 Bt1Gc1281	
	78 100	1.20E-200	1942 1	1991	g1709059	150-1565	81 Bt1G1405	949 Bt1Gc1281	
0 (Z99108) transcriptional regulator [Bacillus subtilis]	50 100	4.60E-162	1578 4	1483	g2633134	417-2277	82 Bt1G1404	948 Bt1Gc1282	
,	81 63	1.70E-22	261	260	g134408	1690-1502	75 Bt1G1403	947 Bt1Gc1275	
M) [Bacillus subtilis] 0 ADENYLATE KINASE (ATP-AMP	75 100	4.90E-87	870	866	g125158	1439-789	75 Bt1G1402	947 Bt1Gc1275	
	80 100	6.30E-110	1086 6	1086	·g113739	789-46	75 Bt1G1401	947 Bt1Gc1275	
subtilis HYPOT	46 70	1.70E-54	563	516	g1176994	2977-2306	80 Bt1G1400	946 Bt1Gc1280	
	66 100	2.20E-178	1732 2	1853	g2828494	1854-210	80 Bt1G1399	946 Bt1Gc1280	
_	49 45	1.40E-41	441	350	g3256407	2577-3092	74 Bt1G1398	945 Bt1Gc1274	
O (AJ005255) OxyR [Erwinia chrysanthemi]	24 100	1.00E-22	263	258	g4583559	2443-1528	74 Bt1G1397	945 Bt1Gc1274	
	٠.،		-						
- Bacillus subfilis [Bacillus subtilis] 6 DNA-DIRECTED RNA POLYMERASE BETA' CHAIN	77 .36	2.00E-168	1638 2	1575	g2507344	1379-88	79 Bt1G1396	. 944 Bt1Gc1279	
	60 100	6.90E-175	1699 6	1746	g98292	2356-671	71 Bt1G1395	943 Bt1Gc1271	
HYPOTHETICAL II.8 KD PROTEIN IN DNAC-RECK INTERGENIC REGION [Bacillus subtilis]	. 00	3.00E-32	353	443	g1413/2	042-322	/1 Bt1G1394	943 Bt1Gc12/1	
recM p		4.00E-44	465	449	g98442	301-1			
					1				
(Z9702		2.30E-57		574	g2224770	2233-2660			
(Z9702		8.40E-12	160	184	g2224768	1388-1184			
9 (Z97025) vlaH [Bacillus subtilis]	57 99	1.20E-28	319	312	£2224767	547-861	77 Bt1G1390	942 Bt1Gc1277	
	% % Ident Cvrg	BlastP- % Prob Ider	BlastP I Score	nap (NCBI gi	Position	ld Gene Id	ID Contig Id	
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[Bacillus Subtilis] (Z92952) unknown [Bacillus subtilis]	31 65	1.60E-08	129 1.0	87	g1894749	2549-2858	0 Bt1Gc1295 Bt1G1429	960
	45 29	1.30E-35	388 1.3	335	g1894748	1-2553	0 Bt1Gc1295 Bt1G1428	960
	29 100	1.50E-12	167 1.5	144	g3258025	2964-3704	9 Bt1Gc1294 Bt1G1427	959
	32 48	1.90E-06	112 1.9	118	g3258025	1424-1741	9 Bt1Gc1294 Bt1G1426	959
_	51 48	4.80E-25	285 4.8	241 . :	g2443233	3699-3373	Bt1Gc1289	958
					(•		
	28 100	1.40E-32	356 1.4	282	g2126561	3092-2208	8 Bt1Gc1289 Bt1G1424	958
NTERGENIC REGION [Saccharomyces cerevisiae]	30 99	4./UE-18 .	219 4.		8/31703	1001-0017	0 DITUCIZOS DITUTAZO	. 930
					7210/2		7176	2
	39 78	4.00E-60	616 4.0	806	g1723606	1343-1	8 Bt1Gc1289 Bt1G1422	958
INTERGENIC REGION [Bacillus subtilis]								
HYPOTHETICAL 137.4 KD PROTEIN IN BCSA-DEGR	35 41	8.70E-81	815 8.7		g1730891	862-2345	7 Bt1Gc1292 Bt1G1421	957
_		9.00E-24	273 9.0		g4894302	300-1	7 Bt1Gc1292 Bt1G1420	957
(U57060) ScdA [Staphylococcus aureus]	34 .41	1.80E-11	157 1.8		g1575061	3083-3367	6 Bt1Gc1291 Bt1G1419	956
_	33 100	1.90E-37	402 1.9	362 4	g1945654	325-1125	6 Bt1Gc1291 Bt1G1418	956
[Bacillus subtilis]								
(SUPEROXIDE-INDUCIBLE PROTEIN 9) (SOI9)								
INDEPENDENT METHIONINE SYNTHASE)								
INDEPENDENT ISOZYME) (COBALAMIN-								
(METHIONINE SYNTHASE, VITAMIN-B12		-						
	51 66	2.60E-136	1335 2.60	1317 13	g3334472	1203-2724	5 Bt1Gc1290 Bt1G1417	955
(U8259	28 . 58	3.70E-20	242 3.7		g1778501	3313-1189	4 Bt1Gc1288 Bt1G1416	954
_	39 80	3.40E-40	428 3.4	358 ′	g1894745	669-1	4 Bt1Gc1288 Bt1G1415	954
(Z75208) hypothetical protein [Bacillus subtilis]	76 100	2.20E-59	609 2.2	881	g1770004	2187-1494	3 Bt1Gc1284 Bt1G1414	953
INFLUENZAE. [Bacillus subtilis]								
PRODUCTS IN SYNECHOCYSTIS AND H.					٠			
(AB001488) FUNCTION UNKNOWN, SIMILAR	50 100	3.30E-58 :	598 3.3	921 5	g1881258	1173-121	3 Bt1Gc1284 Bt1G1413	953
(Y1408	79 100	2.40E-124 (1222 2.40	1624 13	g2226151	771-1994	2 Bt1Gc1287 Bt1G1412	952
(Y14080) hypothetical protein [Bacillus subtilis]	61 99	1.60E-47	497 1.6	532 4	g2226150	281-772	2 Bt1Gc1287 Bt1G1411	952
•	nt Cyrg	Prob Ident		Score Score		•		NO !
NCBI gi description				aat_ BlastP	NCBI gi	Position	Contig Id Gene Id	SEQ

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CARBOXYVINYLTRANSFERASE (ENOYLPYRUVATE TRANSFERASE) (UDP-N-ACETYLGLUCOSAMINE								• ,		
PROBABLE UDP-N-ACETYLGLUCOSAMINE 1-	36		2.80E-59	608	599	g1171068	1-468	Bt1G1451	969 Bt1Gc1304	
INTERGENIC REGION (O357) [Escherichia coli] (AF002191) YhzA homolog [Bacillus subtilis]	99	32	2.40E-23	269	211	£2828809	1482-983	Bt1G1450	968 Bt1Gc1305	
HYPOTHETICAL 39.8 KD PROTEIN IN OSMY-DEOC	88		1.90E-23	270	131	g732119	1284-1	Bt1G1449	968 Bt1Gc1305	
ERYTHROMYCIN ESTERASE TYPE II [Escherichia coli]	32	30	6.20E-19	232	156	g119547	2268-1862	Bt1G1448	967 Bt1Gc1303	
(D86417) YfIK [Bacillus subtilis]	32	49	9.10E-15	188	169	g2443233	3457-3670	Bt1G1447	966 Bt1Gc1293	
colij			•			!				
	100		2.90E-41	438	398	g1657534	3385-2391	Bt1G1446	966.Bt1Gc1293	
COTJA PROTEIN [Bacillus subtilis]	99	-	4.30E-24	276	239	g2833394	2526-2278	Bt1G1445	965 Bt1Gc1301	
COTJB PROTEIN [Bacillus subtilis]	99	52	1.40E-16	205	263	g2833395	2331-2025	Bt1G1444	965 Bt1Gc1301	
COTJC PROTEIN [Bacillus subtilis]	99	89	3.70E-89	890	890	g2833396	2009-1443	Bt1G1443	965 Bt1Gc1301	
subtilis]										
(AF008220) branch-chain amino acid transporter [Bacillus	93	4	5.90E-75	756	859	g2293322	1222-1	Bt1G1442	965 Bt1Gc1301	
(Z99108) similar to hypothetical proteins [Bacillus subtilis]	99	32	1.10E-27	310	277	g2633172	1512-943	Bt1G1441	964 Bt1Gc1296	
STAGE V SPORULATION PROTEIN R [Bacillus subtilis]	57	79	6.70E-106	849	1096	g586024	819-1	Bt1G1440	964 Bt1Gc1296	
(Z99107) similar to hypothetical proteins [Bacillus subtilis]	100		2.90E-98	976	1040	g2632998	2331-1615	Bt1G1439	963 Bt1Gc1299	
(AL049573) MutT-like protein [Streptomyces coelicolor]	89	31	1.60E-10	148	100	g4582371	1414-28	Bt1G1438	963 Bt1Gc1299	
aeolicus]									٠	
(AE000699) penicillin binding protein 1A [Aquifex	16	53	7.70E-31	348	296	g2983239	1541-1	Bt1G1437	963 Bt1Gc1299	
(297112) distrible gene name: ymxi, sminar to deasstylass	70	5	4.70E-07	0/0	037	2404042	7474-1000	00710110	202 1011001420	
[Bacillus subtilis]	0		4 DOE 97	870	830	CV0VE9C	2204-1255		067 B+1C-1708	
HYPOTHETICAL PROCESSING PROTEASE (ORFP)	100	74	1.70E-164	1601	1601	g1176567	1268-42	Bt1G1435	962 Bt1Gc1298	
INTERGENIC REGION [Bacillus subtilis]	•	•		٠	-					
[Arabidopsis thaliana] HYPOTHETICAL 16.6 KD PROTEIN IN COTF-TETB	99	56	3.30E-42	447	433	g586830	3749-4189	Bt1G1434	961 Bt1Gc1297	
(AC006200) putative ribose 5-phosphate isomerase	100	39	3.20E-35	381	382	g4262236	1900-3641	Bt1G1433	961 Bt1Gc1297	
PROTEASE) (PROTEASE TI) [Bacillus subtilis]										
SUBUNIT (ENDOPEPTIDASE CLP) (CASEINOLYTIC				,						
ATP-DEPENDENT CLP PROTEASE PROTEOLYTIC	99	2	1.30E-56	583	637	g3287962	1796-1205	Bt1G1432	961 Bt1Gc1297	
DET CINE BETAINE-BINDING FROTEIN FRECONSON	S	U	1.705-50	373	ŽI.C	811/1721	1647-770		701 011001277	
[Bacillus subtilis]	۲.		1 705 36	303	3	~1171021	622-2401	B+1G1/31	061 B+1C+1207	
	ď	!	!		Score				NO	
NCBI gi description	% ! !	% % Ident Cyro	BlastP-	BlastP Score	aat_	NCBI gi	Position	Gene Id	ID Contig Id	
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						Table 1		ξο.	
Contig Id Gene Id		Position	NCBI gi	aat_ nap Score	BlastP Score	BlastP- Prob	- % % Ident Cvrg	% Cvrg	NCBI gi description
		•					•		ENOLPYRUVYL TRANSFERASE) (EPT) [Bacillus
9 Bt1Gc1304 Bt1G1452		588-1553 g418587	g418587	1414	1427	1414 1427 4.60E-146	86	100	subtilis] 46 86 100 HYPOTHETICAL 34.0 KD PROTEIN IN RHO-MURA
			1	٠,				-	INTERGENIC REGION (ORFQ) [Bacillus subtilis]
9 Bt1Gc1304 Bt1G1453 1823-3102 g143434	1453 1	823-3102	g143434	1802	1825	1802 1825 3.10E-188	84	100	(M97678) Rho Factor [Bacillus subtilis]

NO SEQ

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•	976 Bt1Gc1314 Bt1G1471	976 Bt1Gc1314 Bt1G1470	975 Bt1Gc1311 Bt1G1469	975 Bt1Gc1311 Bt1G1468	974 Bt1Gc1310 Bt1G1467	974 Bt1Gc1310 Bt1G1466	974 Bt1Gc1310 Bt1G1465	974 Bt1Gc1310 Bt1G1464		974 Bt1Gc1310 Bt1G1463	973 Bt1Gc1309 Bt1G1462		973 Bt1Gc1309 Bt1G1461	972 Bt1Gc1308 Bt1G1460	972 Bt1Gc1308 Bt1G1459	971 Bt1Gc1307 Bt1G1458	און שניסקיטער שניסודטר	D+1C21307	970 Bt1Gc1306 Bt1G1456		970 Bt1Gc1306 Bt1G1455	970 Bt1Gc1306 Bt1G1454	969 Bt1Gc1304 Bt1G1453		969 Bt1Gc1304 Bt1G1452
	2268-1634	1-1563	2060-1767	1-1361	3248-1609	2752-706	2251-1745	1736-910		1612-83	1606-647		263-1	2185-3107	1-1093	2711-2540	#J0-03-	727 6636	2923-1589		1139-1519	930-48	1823-3102		588-1553
	g2226252	g3323079	g4894353	g2829802	g2766194	g4894577	g4337125	g2766195		g1075130	g1708641		g2829805	g3080755	g2634337	g2633810	82000011	~2622011	g2635594		g1731010	g2293237	g143434	(g418587
	471	518	218	599	175	86	99	239	٠.	302	1018		167	149	1017	110	1401	1041	1350		103	356	1802	٠,	1414
	458	258	194	380	231	159	148	211		144	1035		186	193	912	126	1007		1307		99	365	1825		1427
	2.20E-43	1.40E-19	2.10E-15	2.70E-65	4.50E-19	4.20E-11	1.60E-10	3.30E-17		1.00E-15	1.60E-104		1.50E-14	3.10E-15	1.70E-91	1.30E-07	/./0E-100	7 70E 160	2.40E-133		2.50E-05	1.60E-33	3.10E-188		4.60E-146
	46	30	48	34	4	27	26	25		27	62		41	25	54	47	8	5	55		31	50	%		86
	2	90	99	100	28	52	99	100		100	100		, 32	%	<u>&</u>	19.	5		100		99	99	100		100
	protein [Treponema pallidum] (Y14084) hypothetical protein [Bacillus subtilis]	(AE001248) conserved hypothetical integral membrane	(AF065404) pXO1-138 [Bacillus anthracis]	hyodysenteriae] HYPOTHETICAL 53.3 KD PROTEIN IN SFP-GERKA	chaffeensis] (U75349) putative ABC transporter BhiD [Brachyspira	[Brachyspira hyodysenteriae] (AF117273) iron-binding protein precursor [Ehrlichia	hyodysenteriae] (U75349) periplasmic-iron-binding protein BhiA	(U75349) putative permease BhiE [Brachyspira	(sfuB) homolog - Haemophilus influenzae (strain Rd KW20) [Haemonhilus influenzae Rd]	periplasmic-binding-protein-dependent iron transport protein	(U51115) YeaC. [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 30.5 KD PROTEIN IN GABP-GUAA	(AF016483) APH(2")-Id [Enterococcus casseliflavus]	(Z99114) similar to hypothetical proteins [Bacillus subtilis]	(Z99111) fructose-1-phosphate kinase [Bacillus subtilis]	(25)111) phosphodalisticase system (r 13) nuclose-specific enzyme [IRC component [Bacillus subtilis]	[Bacillus subtilis]	(Z99119) similar to Na+-transporting ATP synthase	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 13.9 KD PROTEIN IN CCCA-SODA	(AF008220) Yell [Bacillus subtilis]	(M97678) Rho Factor [Bacillus subtilis]		HYPOTHETICAL 34.0 KD PROTEIN IN RHO-MURA

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984 985 985 986	984 984 984	982 982 983	980 981 981 981	979 979 980	978 978	978	SEQ ID NO 977
Bt1Gc1319 Bt1Gc1322 Bt1Gc1322 Bt1Gc1325	Bt1Gc1319 Bt1Gc1319 Bt1Gc1319	Bt1Gc1318 Bt1Gc1318 Bt1Gc1317	Bt1Gc1315 Bt1Gc1312 Bt1Gc1312 Bt1Gc1312	Bt1Gc1302 Bt1Gc1302 Bt1Gc1315	Bt1Gc1316 Bt1Gc1316	978 Bt1Gc1316	Contig Id Bt1Gc1313
Bt1G1492 Bt1G1493 Bt1G1494 Bt1G1495	Bt1G1489 Bt1G1490 Bt1G1491	Bt1G1485 Bt1G1486 Bt1G1488	Bt1G1481 Bt1G1482 Bt1G1483 Bt1G1484	Bt1G1478 Bt1G1479 Bt1G1480	Bt1G1476 Bt1G1477	Bt1G1473 Bt1G1474	Gene Id Bt1G1472
3766-4160 236-679 2601-1080 2757-2050	1-493 515-1135 1149-2097	397-1 2326-532 2455-2102	2345-1881 382-1809 2304-1909 2555-2614	1-573 614-3114 834-1815	1777-2442 1777-2604	214-546 549-1352	Position 1916-387
g141186 g1303698 g953179 g399406	g1204036 g2498126 g4154878	g3258251 g2632414 g2497392	g1749770 g2982907 g2635672 g2127280	g2612898 g1001551 g1703442	g1176704 g1176704 g1730929	g1176702 g1706300	NCBI gi g2633324
187 255 376 246	201 257 504	174 85 584	308 1090 267 108	516 57 779	4/5 668 119	901	nap Score
180 278 445 212	217 186 277	201 145 585	351 1055 273 108	522 202 844	684	-	BlastP Score 594
6.40E-14 2.60E-24 5.30E-42 2.60E-17	7.70E-18 1.50E-14 6.80E-41	3.80E-16 6.70E-10 7.80E-57	4.90E-32 1.20E-106 9.00E-24 1.80E-05	3.70E-50 1.60E-15 2.80E-84	2.10E-47 2.50E-67 2.80E-08	8.40E-44 1.40E-87	BlastP- Prob 8.60E-58
36 43 37 25	29 40	34 30 96	46 39 100		60		% Ident
74 99 100 85	58 100 100	48 46 47	52 100 99 5	59 57 100	100	100	% Cvrg
		(AP000007) 284aa long hypothetical cobalt transport ATP-binding protein [Pyrococcus horikoshii] (Z99104) ybaF [Bacillus subtilis] (INSERTION SEQUENCE IS232 PUTATIVE ATP-BINDING PROTEIN [Insertion sequence IS232]			MEIHYLGLYOXAL SYNIHASE [Bacillus subtilis] HYPOTHETIÇAL 24.8 KD PROTEIN IN DAPB-PAPS INTERGENIC REGION [Bacillus subtilis] HYPOTHETICAL 42.0 KD PROTEIN IN DAPB-PAPS		_

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993 1	993 I	993 1	993 I	993 I	993 I	992 I		992 I	992 I	•	991 E	991 E		991 E		990 I	990 E		989 E			988 E	988 E		988 E		988 E	.988 E	987 E		SEQ
Bt1Gc1329 Bt1G1516	Bt1Gc1329	Bt1Gc1329	Bt1Gc1329	Bt1Gc1329	Bt1Gc1329	Bt1Gc1330		Bt1Gc1330	Bt1Gc1330		Bt1Gc1327	Bt1Gc1327		Bt1Gc1327		Bt1Gc1324	Bt1Gc1324		Bt1Gc1321		-	Bt1Gc1326	Bt1Gc1326	•	Bt1Gc1326	-	Bt1Gc1326	Bt1Gc1326	Bt1Gc1323	Contig Id	
Bt1G1516	Bt1G1515	Bt1G1514	Bt1G1513	Bt1G1512	Bt1G1511	Bt1G1510		Bt1G1509	Bt1G1508		Bt1G1507	Bt1G1506		Bt1G1505		Bt1G1504	Bt1G1503		Bt1G1502		1	Bt1G1501	Bt1G1500	•	Bt1G1499		Bt1G1498	Bt1G1497	Bt1G1496	Gene Id	
4532-3635	3488-2337	2124-1438	1346-1	870-424	248-1	3131-2570		785-1513	1-1319		2273-2177	1795-2273		1656-1		3122-2445	342-1		2698-1			4356-3680	3536-2177		3526-2140		1447-2142	2119-287	1687-1	Position	
g1177018	g462590	g140681	g3915359	g2226251	g2634266	g2632703	-	g3123142	g1934619		g2497392	g1673402		g1321625	(g2649101	g2633840		g3025120	,		e1881353	g2688416		g2314344	(£2633027	g2196513	g1708794	NCBI gi	
880	604	663	112	389	198	293		200	264		148	214		2084		444	314		969		ļ	262	175		620		293	169	2457	nap Score	aat
867	364	672	182	398	3 210	334		246	349		£ 42	304		2064		423	347		925			309	182		493		297	358	2314	Score	Riggi
1.00E-86	3.00E-54	4.70E-66	1.40E-13	5.10E-37	4.20E-17	3.10E-30		6.50E-21	7.90E-32		3.20E-12	9.30E-27		1.50E-213		1.10E-39	1.30E-31		7.30E-93			1.40E-27	2.60E-11		4.40E-47		2.60E-26	8.80E-33	4.70E-240	Prob	Riger P_
60	36	55	26		52	39		27	32		97	42		67		40	58		37		ţ	29	21		32		35	35	85	=	%
2	100	100	<u>4</u> 1	99	51	99.		100	93		13	29		99		100	37		93		;	∞ .	100		100		100	70	92	Cyrg	\$
(MEDH) [Bacillus sp.] HYPOTHETICAL 48.3 KD PROTEIN IN KATB	INTERGENIC REGION [Escherichia coli] NAD-DEPENDENT METHANOL DEHYDROGENASE	fulgidus] HYPOTHETICAL 25.2 KD PROTEIN IN LYSR-ARAE	HYPOTHETICAL PROTEIN AF0433 [Archaeoglobus	(Y14084) hypothetical protein [Bacillus subtilis]	(Z99114) yoaS [Bacillus subtilis]	(Z99106) yczH [Bacillus subtilis]	INTERGENIC REGION [Escherichia coli]	HYPOTHETICAL 26.1 KD PROTEIN IN PROX-MPRA.	(U93874) YrhO [Bacillus subtilis]	BINDING PROTEIN [Insertion sequence IS232]	INSERTION SEQUENCE IS232 PUTATIVE ATP-	(Z82044) hypothetical 54.4 kd protein [Bacillus subtilis]	stearothermophilus]	(D84648) exo-alpha-1,4-glucosidase [Bacillus	[Archaeoglobus fulgidus]	(AE001001) ABC transporter, ATP-binding protein	(Z99111) yktD [Bacillus subtilis]	PROTEIN YDIF [Bacillus subtilis]	HYPOTHETICAL ABC TRANSPORTER ATP-BINDING	REGULATORS IN B.SUBTILIS. [Bacillus subtilis]	SIMILAR TO MULTIDRUG-EFFLUX TRANSPORTER	protein [Borrella burgdorteri] (AB001488) PROBABLE REGULATORY PROTEIN.		protein [Helicobacter pylori 26695]	(AE000624) conserved hypothetical integral membrane	[Bacillus subtilis]	epidermidis] (Z99107) similar to hypothetical proteins from B. subtilis	(U77778) putative membrane protein [Staphylococcus	GTP-BINDING PROTEIN LEPA [Bacillus subtilis]	NCBI gi description	

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1001 Bt1Gc1336 Bt1G1535	1001 Bt1Gc1336 Bt1G1534	1000 Bt1Gc1338 Bt1G1533	1000 Bt1Gc1338 Bt1G1532	21.00	1000 Bt1Gc1338 Bt1G1531	1000 Bt1Gc1338 Bt1G1530	999 Bt1Gc1337 Bt1G1529		998 Bt1Gc1334 Bt1G1528	998 Bt1Gc1334 Bt1G1527	998 Bt1Gc1334 Bt1G1526		998 Bt1Gc1334 Bt1G1525	998 Bt1Gc1334 Bt1G1524	997 Bt1Gc1335 Bt1G1523		996 Bt1Gc1333 Bt1G1522		995 Bt1Gc1332 Bt1G1521		1	995 Bt1Gc1332 Bt1G1520		994 Bt1Gc1331 Bt1G1519		994 Bt1Gc1331 Bt1G1518	994 BtlGcl331 BtlGl517		SEQ ID Contig Id Gene Id NO
2686-1694	1707-907	3373-4292	3233-1844		3071_1803	1307-456	1283-150		4968-4643	4456-3720	3686-2862		2842-2408	1231-2361	2008-200		304-1008		3494-97			1618-1		2129-2659	1	588-2111	35-481	•	Position
g2126200	g1706796	g1176933	g2293169		03756503	g1864009	g1926326		g1731008	g2415737	g3758894		g1730119	g1730252	g1865711		g3608398		g2388582		0	g1174516		g1170427	0	0123758	g123762		NCBI gi
545	569	190	113		453	692	693		442	403	356		496	1640	458		446		122			1843		721	. !	1825	442	:	nap Score
515	587	336	162	,	ŝ	705	724		458	480	373		472	1579	597		482		197	•		1846	Ą	646	:	1777	462	;	BlastP Score
2.00E-49	4.80E-57	1.90E-30	5.20E-12	0.000	3 00E-48	··· 1.50E-69	1.40E-71		2.20E-43	1.00E-45	2.30E-34		7.30E-45	3.60E-162	4.20E-58		6.40E-46		1.60E-14			1.80E-190		2.70E-63		3 80E-183	8.40E-44		BlastP- Prob
38	42				٠ ع	49	39		81		31		63		35		46		27			58		77		70	5		% % Ident Cyrg
100	100	,100	&	5	<u> </u>	100	100		37	100	100		99	100	100		100		27		;	<u> </u>		32		1	99	3	% Cvrg
hemin permease - Yersinia enterocolitica []	S'REGION [Buchnera aphidicola] FERRICHROME TRANSPORT ATP-BINDING PROTEIN FHIIC [Bacillus subtilis]	HYPOTHETICAL 34.4 KD PROTEIN IN TRPA	(AF008220) transcription regulator [Bacillus subtilis]	horikoshiil	purpuras [Escherichia coli] A D0000013 42822 long hypothetical protein [Pyrococcus	(D85892) homlogue of rdmC protein of Streptomyces	(X98106) integrase [Bacteriophage phig1e]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 32.5 KD PROTEIN IN CCCA-SODA	(AB000617) YcdI [Bacillus subtilis]	(Z71552) AdcB protein [Streptococcus pneumoniae]	1 [Bacillus subtilis]	FERRIC UPTAKE REGULATION PROTEIN HOMOLOG	GCPE PROTEIN HOMOLOG [Bacillus subtilis]	(Y11477) endolysin [Bacteriophage Bastille]	faecalis]	(AF071085) putative glycosyl transferase [Enterococcus	transferase (gb U76557). [Arabidopsis thaliana]	aureus] (AC000098) Contains similarity to Rattus O-GlcNAc	(MUPIROCIN RESISTANCE PROTEIN) [Staphylococcus	RESISTANT (ISOLEUCINETRNA LIGASE) (ILERS)	ISOLEHCYL-TRNA SYNTHETASE MUDIROCIN	(IMIDAZOLONEPROPIONATE HYDROLASE) [Bacillus	UROCANATE HYDRATASE (UROCANASE)	subtilis	HISTIDINE AMMONIA-I VASE (HISTIDASE) [Bacillus	HUT OPERON POSITIVE REGULATORY PROTEIN	3'REGION [Bacillus subtilis]	NCBI gi description

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[Synechocystis sp.]								-		
typhimurium] HYPOTHETICAL 25.5 KD PROTEIN SLR0014	43 100		2.40E-46	486	517	g2833487	520-1215	0 Bt1G1563	1010 Bt1Gc1350	
family) [Bacillus subtilis] MG(2+) TRANSPORT ATPASE, P-TYPE 2 [Salmonella	65 16		8.10E-50	526	526	g1168550	1-444	0 Bi1G1562	1010 Bt1Gc1350	
[Bacillus subtilis] (Z99119) similar to transcriptional regulator (AraC/XylS	29 22	•	1.50E-14	197	159	g2635499	1-2439	9 Bt1G1561	1009 Bt1Gc1349	
(Z83337) similar to B. subtilis YcsE hypothetical protein	32 100		7.00E-40	425	393	g1763711	291-1136	7 Bt1G1560	1008 Bt1Gc1347	
SENSOR PROTEIN RESE [Bacillus subtilis]		_	4.70E-137	1342	1491	g466195	2578-806			
(Z99120) similar to hypothetical proteins [Bacillus subtilis]			2.80E-59	608	597	g2635812	662-87			
hypothetical protein - Escherichia coli [Escherichia coli]	49 99		9.00E-24	273	371	g421055	4805-4367	5 Bt1G1557	1006 Bt1Gc1345	
COLLAGENASE) [Clostridium perfringens]	· • .									
MICROBIAL COLLAGENASE PRECURSOR (120 KD	47 88		1.80E-240	2318	2254	g1169000	3920-1017	5 Bt1G1556	1006 Bt1Gc1345	
kinB protein - Bacillus subtilis []	34 50		3.40E-25	287	312	g421510	1-646	5 Bt1G1555	1006 Bt1Gc1345	
fulgidus]										
(AE001068) conserved hypothetical protein [Archaeoglobus	35 . 33		4.30E-07	121	122	g2650107	2796-3085	4 Bt1G1554	1005 Bt1Gc1344	
[Pseudomonas putida]										
BKD OPERON TRANSCRIPTIONAL REGULATOR	37 99		8.00E-23	264	270	g1168677	2575-2096	4 Bt1G1553	1005 Bt1Gc1344	
(AE000736) hypothetical protein [Aquifex aeolicus]	41 99		8.40E-28	311	304	g2983780	2006-1474	4 Bt1G1552	1005 Bt1Gc1344	
tuberculosis]						•				
(Z74020) hypothetical protein Rv1544 [Mycobacterium	33 77		9.50E-20	235	150	g1403498	638-2439	4 Bt1G1551	1005 Bt1Gc1344	
protein (MerR) [Listeria monocytogenes]										
(Y07640) putative mercury resistance operon regulatory	37 45		3.30E-10	145	110	·g2879772	531-335	4 Bt1G1550	1005 Bt1Gc1344	
(AB013368) YkfB [Bacillus halodurans]	50 97		1.20E-69	706	756	g4514327	2820-3798	3 Bt1G1549	1004 Bt1Gc1343	
INTERGENIC REGION (ORFC) [Bacillus subtilis]									5-	
HYPOTHETICAL OXIDOREDUCTASE IN NATB-RAPJ	36 100	•	4.60E-52	540	445	g3183577	1295-363	3 Bt1G1548	1004 Bt1Gc1343	
(AJ237976) hypothetical protein [Streptomyces coelicolor]	37 48		8.40E-12	160	160	g4585854	14-1712	3 Bt1G1547	1004 Bt1Gc1343	
subtilis]	,				-					
ATP-DEPENDENT DNA HELICASE RECG [Bacillus	76 57	-	7.50E-146	1425	1516	g3914611	2488-1315	2 Bt1G1546	1003 Bt1Gc1342	
(Y13937) YlpC protein [Bacillus subtilis]	73 99	-	9.40E-68	688	685	g2337817	1217-654	2 Bt1G1545	1003 Bt1Gc1342	
(Y13937) putative PlsX protein [Bacillus subtilis]	69 64	-	6.50E-76	765	742	g2337818	633-1	2 Bt1G1544	1003 Bt1Gc1342	
HI1721 [Haemophilus influenzae Rd]					٠.					
HYPOTHETICAL TRANSPOSASE-LIKE PROTEIN	40 57	_	2.50E-19	231	183	g2497400	1768-1415		1002 Bt1Gc1341	
COMA OPERON PROTEIN 2 [Bacillus subtilis]	57 99		3.20E-35	381	380	g116904	962-1339	1 Bt1G1537	1002 Bt1Gc1341	
(AF015825) NodB-like protein [Bacillus subtilis]	7 48		1.20E-13	185	104	g2612882	1768-1	1 Bt1G1536	1002 Bt1Gc1341	
	11 (18	TOCHL	1100	91036	·	•			NO	
NCBI gi description		I %	BlastP-	BlastP	nap _	NCBI gi	Position	Gene Id	ID Contig Id	
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1019 Bt1Gc1359	1018 Bt1Gc1348	1018 Bt1Gc1348	1017 Bt1Gc1358			1015 Bt1Gc1356		1015 Bt1Gc1356		1015 Bt1Gc1356		1014 Bt1Gc1353		1014 Rt1Gc1353		1014 B+1C+1353		1014 Bt1Gc1353		1014 Bt1Gc1353			1013 Bt1Gc1355	1012 Bt1Gc1352	1012 Bt1Gc1352			1011 Bt1Gc1351		1010 Bt1Gc1350	NO Config to	
9 Bt1G1583	8 Bt1G1582	8 Bt1G1581	8 Bt1G1580	8 Bt1G1579	7 Bt1G1578	6 Bt1G1577		6 Bt1G1576		6 Bt1G1575		3 Br1G1574		3 R+1G1573		3 B+1G1573		3 Bt1G1571		3 Bt1G1570			5 Bt1G1568	2 Bt1G1567	2 Bt1G1566			1 Bt1G1565		0 Bt1G1564	Gene Id	
682-1	2433-1479	1393-1	2516-1638	1355-786	1840-993	3298-2374		1310-2007		327-1302		3135-2962		3135_2312	2141-2422	2240-1410	,	441-1355		139-1		3711-2250	1258-1	4800-3647	2586-3611			1335-258	٠	2488-1247	rosition	
g1731364	g2689897	g1168885	g98344	g2635871	g2632519	g1934656		g2829657	C	e3123125	0.10.00	۶1731091	0.00	01731087	84000000	27625852	9	g1731092	•	g1731093	9	92811053	g2632729	g1651216	g4378847			g3023206		g1731071	NCBI gi	
340	667	788	465	258	1138	793		170		714		= .`	Ç	367	.0/1	974	ò	950		117	ۇڭ.	70	616	949	370			606		1108	nap Score	aat
425	721	784	386	270	1151	653		179		576	8	6	į	445	07/	807	ò	963		133	;	11	686	899	482			667		1080	Score	BlastP
7.00E-40	3.00E-71	6.30E-78	9.50E-36	1.90E-23	8.20E-117	4.80E-64		8.20E-14	;	7.00E-56	0.010	0 046	0.000	5 30F-40	0.70E*90	6 705 00		6.80F-97		6.10E-09		8 30F-06	1.50E-67	4.10E-90	6.40E-46			1.60E-65		2.70E-109	Prob	BlastP-
39	45	37	34	31	71	50		25	i	45	Ş	<u>بر</u>	Ų	۵ م	2	5	ç	60		55	ţ	36	35	47	30			45		51	=	%
55	. 89	72	. 100	99	88	100		100		100		77		3 .	100			90		25			•	61		=		100		100	Cvrg	%
HYPOTHETICAL PROTEASE IN ROCR-PURA INTERGENIC REGION [Bacillus subtilis]	subtuis] (AE000792) outer surface protein, putative [Borrelia burgdorferi]	(fragment) [Bacillus subtilis] PUTATIVE CEL OPERON REGULATOR [Bacillus	hypothetical protein (comG 5' region) - Bacillus subtilis	(Z99121) yvaF [Bacillus subtilis]	(Z99105) similar to hypothetical proteins [Bacillus subtilis]	(U93876) cation transport protein YrdO [Bacillus subtilis]	IN DCP-NOHA INTERGENIC REGION [Escherichia coli]	HYPOTHETICAL TRANSCRIPTIONAL REGULATOR	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 35.3 KD PROTEIN IN CSPC-NAP	INTERCENTE DECION [Decilles miletical	HYPOTHETICAL 9 KD PROTEIN IN GI NO-ANSR	INTERCENT DECION [Decilles substite]	HYDOTHETICAL 30 0 KD DROTEIN IN GLNO-ANGR	metabolite debudrorenese [Bacillus subtilis]	(700121) oftemate came name: units such its to affect	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL OXIDOREDITICTASE IN ANSR-BMRIT	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 21.0 KD PROTEIN IN GLNQ-ANSR	Laborational	DI-/TRIPEPTIDE TRANSPORTER (Lactobacillus	(Z99106) vdaL [Bacillus subtilis]	(D88209) Pz-peptidase [Bacillus licheniformis]	(AF124349) unknown [Zymomonas mobilis]	[Escherichia coli]	CARBOXYLATE DEAMINASE (ACC DEAMINASE)	PUTATIVE 1-AMINOCYCLOPROPANE-1-	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 47.0 KD PROTEIN IN GLNQ-ANSR	NCBI gl description	

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	24 °10 39 100	4.00E-08 2.00E-67	141 685	44 699	g3549261 g1934780	1-4096 3783-2515	1024 Bt1Gc1368 Bt1G1601 1024 Bt1Gc1368 Bt1G1602	
<pre>INHIBITOR) {Bacillus subtilis} 9 DNA-ENTRY NUCLEASE (COMPETENCE-SPECIFIC</pre>	57 .99	1.20E-42	451	445	g1171856	1500-1063	1024 Bt1Gc1368 Bt1G1600	
COMP	30 99	1.90E-14	185	149	g112708	1044-652	1024 Bt1Gc1368 Bt1G1599	
[Bacillus subtilis]								
(SUPEROXIDE-INDUCIBLE PROTEIN 9) (SOI9)			-					
INDEPENDENT METHIONINE SYNTHASE)	:	•						
INDEPENDENT ISOZYME) (COBALAMIN-		ř						
-HOMOCYSTEINE WEIHYLIRANSFERASE								
	63 26	5.20E-67	681	653	g3334472	1-592	1024 Bt1Gc1368 Bt1G1598	
			}			}		
0 (D88802) S. lividans chloramphenicol resistance protein;	46 100	2.70E-77	778	952	g1945096	2-1581	1023 Bt1Gc1367 Bt1G1597	
2 (AJ002571) YkbA [Bacillus subtilis]	63 22	3.60E-12	171	300	g2632007	1-289	1023 Bt1Gc1367 Bt1G1596	
	50 100	6.50E-76	765	748	g2632226	2558-3415	1022 Bt1Gc1363 Bt1G1595	
INTERGENIC REGION [Bacillus subtilis]						•		
	66 100	6.30E-78	784	872	g3915503	2328-1567	1022 Bt1Gc1363 Bt1G1594	
	55 100	1.20E-113	1121	1196	g2632230	1309-95	1022 Bt1Gc1363 Bt1G1593	
7 (AF015825) unknown [Bacillus subtilis]	79 87	1.40E-25	217	490	g2612900	2249-1883	1021 Bt1Gc1360 Bt1G1592	
subtilis]								
_	85 100	1.00E-157	1537	1700	g2612901	1838-663	1021 Bt1Gc1360 Bt1G1591	-
				٠,	(
8 TYROSYL-TRNA SYNTHETASE (TYROSINETRNA	51 28	3.70E-19	234	255	g135197	1-348	1021 Bt1Gc1360 Bt1G1590	
	26 100	1.30E-23	7/0	700	81001333	2100-1007	וסבס פנוסכוסטו פניסוס87	
(APUZ)		9.10E-80	370	200	g2019032	2106 1887		
			3					
	44 81	2.40E-53	552	528	g3334448	1-713	1020 Bt1Gc1361 Bt1G1587	
lactis]				-				
(U7333	30 80	2.10E-77	779	555	g4098081	3758-1504	1019 Bt1Gc1359 Bt1G1586	
			ļ	į	- 0			
(T)7333		7 30F_29	<u>კ</u>	212	94098082	3097-1510	Rt1Gc1350	
0 (AJ001103) arcA [Lactococcus lactis]	43 100	4.40E-47	493	460	g3236110	1575-895	1019 Bt1Gc1359 Bt1G1584	
			0.0	**			NO	
NCBI gi description	Ident Cyra	Proh Id	Score	•	NCBI gi	Position	ID Contig Id Gene Id	
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(D78508) YfiJ [Bacillus subtilis] (AL049754) putative two-component system regulator	60 100	30 35	1.90E-28 3.10E-30	317 334	238 290	g1817534 g4753877	1-834 738-1462	1031 Bt1Gc1373 Bt1G1620 1031 Bt1Gc1373 Bt1G1621	
HYPOTHETICAL 39.8 KD PROTEIN IN OSMY-DEOC	73	25	2.20E-20	241	112	g732119	3317-4900	Bt1Gc1374	
(Z99108) yfiT [Bacillus subtilis]	8 8	<u>51</u>	1.50E-46 8 30E-37	3 48 8 8	346 384	g2633163	2499-3027	1030 Bt1Gc1374 Bt1G1617	
tuberculosis]					•				
(Z84498) hypothetical protein Rv1937 [Mycobacterium	58	27	2.80E-45	483	346	g1806225	1-2437	1030 Bt1Gc1374 Bt1G1616	
(D64000) hypothetical protein [Synechocystis sp.]	50	35	1.90E-23	270	240	g1001585	1-496	1030 Bt1Gc1374 Bt1G1615	
HOMOLOG [Bacillus subtilis]									
SUBJULIES PROBABLE ATP-DEPENDENT HELICASE DING	99	39	1.60E-184	1790	1653	g1706437	1143-3912	1029 Bt1Gc1372 Bt1G1614	
(ASPARTATE ALPHA-DECARBOXYLASE) [Bacillus									
ASPARTATE 1-DECARBOXYLASE PRECURSOR	99	74	1.30E-45	479	479	g1709570	631-1011	1029 Bt1Gc1372 Bt1G1613	
ACTIVATING ENZYME) [Bacillus subtilis]								•	
(PANTOTHENATE SYNTHETASE) (PANTOATE						(
PANTOATE-BETA-ALANINE LIGASE	73	55	7.50E-59	604	584	g1709569	1-631	1029 Bt1Gc1372 Bt1G1612	
acyl-carrier protein reductase [Bacillus subtilis]	. [4	10-01	04.0	j	8200000	7203-3730	61.001071	
(700110) alternate cane name: vidA: similar to 3-ovosovi-	3 3	40	4 SOE-61	707	7 T	67633557	4209-3436	Bt1Gc1371	
(AB016282) ORF48 [hacterionhage phi-105]	100 110	2 9	2.50E-35	382	243	94126634	411-2310	Bt1Gc1371	
PYRUVATE KINASE (PK) [Thermococcus litoralis]	 	37	3.30E-33	362	344	£4033437	580-1	1028 Bt1Gc1371 Bt1G1609	
GLUTAMINE CHAIN) [Bacillus caldolyticus]								,	
(CARBAMOYL-PHOSPHATE SYNTHETASE									
PYRIMIDINE-SPECIFIC, SMALL CHAIN				,		(
CARBAMOYL-PHOSPHATE SYNTHASE,	100	71	1.70E-141	1384	1364	g1705597	1202-2291	1027 Bt1Gc1369 Bt1G1608	
DIHYDROOROTASE (DHOASE) [Bacillus caldolyticus]	93	71	3.10E-149	· 1457	1506	g1172784	1-1196	1027 Bt1Gc1369 Bt1G1607	
(03433) 3-Keioacyi-acyi carrier protein reductase [Daciilus subtilis]		ŧ	1,306-34	JO#) + 0	81202421	0071-1107	זמקט פוניסבוסט פוניסוממס	
(Alfo27) 2 lettoral coul possits senting submits submits]	3 8	2 6	1 205 54	07/1	1/10	21502421	2017 1202	Del Ge 1365	
[Bacteriophage SPBc2]	83	79	4 10E-177	1720	1716	~2619016	1-1733	1026 B+1G-1365 B+1G1605	
(Z99115) similar to aminoglycoside N3'-acetyltransferase	100	56	1.40E-73	743	. 789	g2634583	3230-2417	1025 Bt1Gc1364 Bt1G1604	
jannaschii]				,	. .		: : :		
FLAVOPROTEIN HOMOLOG [Methanococcus									
[Bacillus subtilis] DNA/PANTOTHENATE METABOLISM	57	29	9.40E-22	256	173	g3023642	3364-1743	1025 Bt1Gc1364 Bt1G1603	
NCBI gi description	Cvrg	% Ident	BlastP- Prob	BlastP Score	nap Score	NCBI gi	Position	ID Contig Id Gene Id	
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[Bacillus subtilis] (Z82044) hypothetical 28.3 kd protein [Bacillus subtilis] DNA-DIRECTED RNA POLYMERASE ALPHA CHAIN	82 27	29 95	4.00E-28 9.20E-38	314 405	199 389	g1673400 g133395	2698-927 1-259	Bt1Gc1384 Bt1G1643 Bt1Gc1381 Bt1G1644	1041 B 1042 B
PROTEIN 2 IN GLYBC 3 REGION [Bacillus subtilis] (Z99113) similar to hypothetical proteins from B. subtilis	100	73	8.50E-154	1500	1758	g2634220	546-1931	Bt1Gc1384 Bt1G1642	1041 B
HYPOTHETICAL ABC TRANSPORTER ATP-BINDING	100	46	6.40E-133	1303	1389	g1723607	299-2116	Bt1Gc1370 Bt1G1641	1040 B
(L36381) orfA; putative [Neisseria gonorrhoeae]	74	28	7.60E-11	151	93	g790865	933-1	Bt1Gc1380 Bt1G1640	1039 B
INTERGENIC REGION (ORFX) [Bacillus subtilis]	٠.					(
INTERGENIC REGION (ORF1) [Bacillus subtilis] HYPOTHETICAL 38.5 KD PROTEIN IN PONA-COTD	100	46	4.80E-73	738	688	g1730944	2590-3543	Bt1Gc1382 Bt1G1639	1038 B
HYPOTHETICAL 24.0 KD PROTEIN IN PONA-COTD	100	63	2.30E-66	675	645	g732244	2432-1815	Bt1Gc1382 Bt1G1638	1038 B
PENICILLIN-BINDING PROTEINS (A/1B (PBP1) Bacillus subtilis	2	20	1.50E-142	1394	1360	g/302/3	1-00-1	BUGG1382 BUG1637	1038 B
(Z83864) menG [Mycobacterium tuberculosis]	99	<u>.</u>	9.40E-29	320	310	g1781103	1152-1622		
INTERGENIC REGION [Bacillus subtilis]									
HYPOTHETICAL 41.5 KD PROTEIN IN AMHX-AMYE	61	66	2.10E-70	713	842	g2495457	724-1	Bt1Gc1383 Bt1G1635	1037 B
(Z99122) ywzB _i [Bacillus subtilis]	42	67	0.094	65	101	g2636203	1774-1678	Bt1Gc1379 Bt1G1634	1036 B
carboxyvinyltransferase [Bacillus subtilis]				•					
(Z81356) UDP-N-acetylglucosamine 1-	67	76	6.80E-113	1114	1143	g1648861	876-1	Bt1Gc1379. Bt1G1633	1036 B
(Z93940) asparagine synthetase [Bacillus subtilis]	100	45	1.50E-149	1460	1444	g1934835	3582-1756	Bt1Gc1377 Bt1G1632	1035 B
(AE000722) hypothetical protein [Aquifex aeolicus]	21	23	2.40E-08	139	42	g2983568	1047-247	Bt1Gc1378 Bt1G1631	1034 B
(Z99117) similar to hypothetical proteins [Bacillus subtilis]	3 8	30	3.30E-07	118	74	g2635194	482-247	Bt1Gc1378 Bt1G1630	1034 B
(Y08559) Unknown [Bacillus subtilis]	86	45	2.70E-47	495	422	g1592705	3095-3814	Bt1Gc1375 Bt1G1629	1033 B
tuberculosis]						•			
(Z96072) hypothetical protein Rv2688c [Mycobacterium	100	37	4.60E-43	455	437	g2181979	1347-2228	Bt1Gc1375 Bt1G1628	1033 B
(Z99105) similar to protein kinase [Bacillus subtilis]	100	33	1.20E-35	385	374	g2632489	3450-2651	Bt1Gc1376 Bt1G1626	1032 B
(D89936) sporulation-control protein (spo0M) [Bacillus subtilis]	100	4.	1.10E-50	527	471	g3522984	2651-1876	Bt1Gc1376 Bt1G1625	1032 B
megaterium]				-					
hypothetical protein 2 - Bacillus megaterium [Bacillus	100	31	9.70E-27	301	290	g322157	1820-1174	Bt1Gc1376 Bt1G1624	1032 B
170) [Bacillus sp.] HYPOTHETICAL 31.8 KD PROTEIN IN GABP-GUAA	100	51	1.70E-61	629	754	g2828522	297-1172	Bt1Gc1376 Bt1G1623	1032 B
[Streptomyces coelicolor] beta-lactamase (EC 3.5.2.6) precursor - Bacillus sp. (strain	100	81	7.80E-128	1255	1257	g80141	3375-2427	Bt1Gc1373 Bt1G1622	1031 B
NCBI gi description	% Cvrg	% Ident	BlastP- Prob	BlastP Score	nap Score	NCBI gi	Position	Contig Id Gene Id	SEQ O

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(U68235) SpoIVB [Bacillus subtilis]	100	42 64	6.30E-142	1388	1365	g1553038	3337-2057	1050 Bt1Gc1393 Bt1G1665	٠
STAGE 0 SPORULATION PROTEIN A [Bacillus	100	34 99	1.30E-134	1319	1303	g1711477	1760-968	1050 Bt1Gc1393 Bt1G1664	_
(H1) [Bacillus subtilis] (AF065404) pXO1-90 [Bacillus anthracis]	37		1.10E-48		428	g4894306	1977-2709	Bt1Gc1392	:
	62	25 35	1.90E-25	227	595	g730002	1-1531	1049 Bt1Gc1392 Bt1G1662	_
(AB001488) SIMILAR TO YDDS_BACSU. [Bacillus	3 43	27 73	4.60E-27	304	686	g1881327	3253-2695	1048 Bt1Gc1391 Bt1G1661	_
(Y14083) Hypothetical protein [Bacillus subtilis]	100		1.40E-105	1045	1119	g2226235	1656-2630	1048 Bt1Gc1391 Bt1G1660	_
aerugınosa] (U66480) YnbA [Bacillus subtilis]	8 6.	61 48	2.20E-61	628	634	g1750108	889-1	1048 Bt1Gc1391 Bt1G1659	
FIMT 3'REGION (DADA*) (ORFZ) [Pseudomonas									
HYPOTHETICAL 39.5 KD OXIDOREDUCTASE IN	73	20 25	5.60E-20	238	157	g3915984	3305-2440	1047 Bt1Gc1387 Bt1G1658	_
(VEGETATIVE PROTEIN 81) (VEG81) [Bacillus subtilis]									
INTERGENIC REGION (O162) [Escherichia coli] DIHYDRODIPICOLINATE SYNTHASE (DHDPS)	48	23 33	1.10E-23	272	248	g416876	421-1	1047 Bt1Gc1387 Bt1G1657	_
HYPOTHETICAL 18.8 KD PROTEIN IN GNTR-GGT	99	09 27	2.90E-09	136	89	g1176283	941-426	1046 Bt1Gc1390 Bt1G1656	_
(Y14081) hypothetical protein [Bacillus subtilis]	22	_	5.60E-17	210	210	g2226185	225-1	1046 Bt1Gc1390 Bt1G1655	_
ELEMENT IS232 [Insertion sequence IS232]					~*				-
TRANSPOSASE FOR INSERTION SEQUENCE	45	88 91	2.60E-88	882	912	g2497382	3617-4206	1045 Bt1Gc1385 Bt1G1654	_
licheniformis						•			
(AF007865) bacitracin synthetase 1; BacA [Bacillus	16	50 38	6.80E-150	1485	1392	g2982194	1-4206	1045 Bt1Gc1385 Bt1G1653	_
(AF082668) CsrR [Streptococcus pyogenes]	100	38 41	2.10E-38	411	432	g3599371	993-313	1044 Bt1Gc1386 Bt1G1652	_
INTERGENIC REGION [Bacillus subtilis]		•			:	. '	٠		
HYPOTHETICAL 30.8 KD PROTEIN IN SINI-GCVT	38		1.90E-23		247	g1731039	3872-3575	1043 Bt1Gc1388 Bt1G1651	Ľ
(Z99116) yqzE [Bacillus subtilis]	99		1.80E-09	138	150	g2634900	1844-2955	1043 Bt1Gc1388 Bt1G1650	_
SHIKIMATE KINASE (SK) [Erwinia chrysanthemi]	99		1.50E-14		187	g114199	1305-1811	1043 Bt1Gc1388 Bt1G1649	_
(AC004450) hypothetical protein [Arabidopsis thaliana]	76		1.70E-29	327	222	g3763917	1-1189	1043 Bt1Gc1388 Bt1G1648	_
(D64126) unknown [Bacillus subtilis]	95	80 61	6.20E-80	803	852	g1644203	1621-2405	1042 Bt1Gc1381 Bt1G1647	_
PROTEIN YBXA [Bacillus subtilis]	-			•					•
subtilis] HYPOTHETICAL ABC TRANSPORTER ATP-BINDING	100	87 60	8.00E-87	868	831	g3915966	761-1603	1042 Bt1Gc1381 Bt1G1646	_
50S RIBOSOMAL PROTEIN L17 (BL15) (BL21) [Bacillus	. 99	51 88	8.50E-51	528	528	g132718	298-657	1042 Bt1Gc1381 Bt1G1645	
(TRANSCRIPTASE ALPHA CHAIN) (RNA POLYMERASE ALPHA SUBUNIT) [Bacillus subtilis] 👶	٠.					٠,			
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)58)57	59	235	5254	1051		1270			1559		<u> </u>		53	Ю35			292			1429		20	314		929)578	49	732	51 <u>81</u>	•
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ENOLPYRUVYLSHIKIMATE-3-PHOSPHATE PHOSPHOLYASE) (VEGETATIVE PRÓTEIN :	CHORISMATE SYNTHASE (5-	IVHB	ORF IS231C [Bacillus thuringiensis]	illine .	(Y14084) hypothetical protein [Bacillus subtilis])0075	[Mycobacterium tuberculosis]	HYPOTHETICAL 41.2 KD PROTEIN CY277.09	(PSEUDOURIDYLATE SYNTHASE) (URACIL	UDO	TRNA PSEUDOURIDINE SYNTHASE B (TRNA	[Bacillus subtilis]	RIBOSOME-BINDING FACTOR A (P15B PROTEIN)	INTERGENIC REGION (ORF5) [Bacillus subtilis]	HYPOTHETICAL 10.7 KD PROTEIN IN INFB-RPSO	(Z99112) initiation factor IF-2 [Bacillus subtilis]	(PHERS) [Bacillus subtilis]	ENYI	IXN.	CHAIN) (PHERS) [Bacillus subtilis]		PHENYLALANYL-TRNA SYNTHETASE ALPHA	ANSC	A-DIR	00822	INTERGENIC REGION [Bacillus subtilis]	HTO	POLY(A) POLYMERASE (PAP) [Bacillus subtilis]	(U20445) BirA protein [Bacillus subtilis]	05599		
PYRUVYLSHIKIMATE-3-PHOSPHATE PHOLYASE) (VEGETATIVE PROTEIN 216)	MATE	$\frac{3}{2}$		ETIC	hypo	8) hy	teriun	ETIC	VASI	URID URID	EUD	subtili	VE-BI	NIC	ETIC	initiat	Baci	ALA	ALA	PHE	HEN	YLALANYL-TRNA SYNTHE	RIPI	ECT1	(0) pu	OINE	ETIC	POL	BirA	5996) RêcN homolog [Bacillus anthracis]	. ,	
YLSH \SE) (SYN	Ĭ	Bacill	AL P	thetica	othet	n tube	AL 4:		NES	OURI	<u>s</u>	NUN	REGI	AL 10	ion fa	illus s	NINE	¥,	S) []	YLA		ASE	ED RI	tative	REGI	ALA:	YME	prote	eN ho	2	` !
[VEG]	AHT	TES	us thu	ROI'E	al pro	ical p	rculo	1.2 KI	TE S	5 SY	DINE		VG F/	9 (2)).7 KI	ctor I	ubtilis	TR	TRNA	Bacill	ANI	TRNA	BET/	VA PO	cyste	28	2.0 KI	RASE	<u>≅</u> ≅.	molo	CBI	
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3-PHO	•	HASE	nsis]	HUA	Bacill	Вас		DIE!	IASE	\SE)	THAS		RA() [Ba	TEI	Bacill		[GAS]	THE	tilis]	RNA	THE		TERA	nthase	us sut	TEI) [Ba	subti	cillus	NCB1 gi description	:
OSPH PROT		Bac		A 5'K	us sub	illus n		CY2	(UR	PSI5	SEB(P15B	illus	I N I	us sut		E BET	FASE	!	/DIT	[ASE	RNA	SE BI	Bac	tilis]		cillus	lis]	anthr	'n	•
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16)		3-DEHYDROQUINATE SYNTHASE [Bacillus subtilis]		HYPOTHETICAL PROTEIN IN PDHA 5'REGION (ORFI)	<u>;</u>	(AJ000758) hypothetical protein [Bacillus megaterium]				PSEUDOURIDINE 55 SYNTHASE) (PSI55 SYNTHASE)	**		EIN)	S	PSO			(PHENYLALANINETRNA LIGASE BETA CHAIN)	PHENYLALANYL-TRNA SYNTHETASE BETA CHAIN	į	CHAIN (PHENYLALANINETRNA LIGASE ALPHA	Š.	(TRANSCRIPTASE BETA CHAIN) (RNA POLYMERASE	DNA-DIRECTED RNA POLYMERASE BETA CHAIN	(AF008220) putative cysteine synthase [Bacillus subtilis]		HYPOTHETICAL A2.0 KD PROTEIN IN DAPB-PAPS	is]				
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	Bt1Gc1406	Bt1Gc1407	Bt1Gc1407	BIIGC140/		Bt1Gc1404	Bt1Gc1404			Bt1Gc1404		Bt1Gc1405		1061 Bt1Gc1405		Bt1Gc1405		Bt1Gc1402			Bt1Gc1402	Bt1Gc1402	Bt1Gc1403	Bt1Gc1403		Bt1Gc1403	Bt10c1403		Bt1Gc1400		1058 Bt1Gc1400		Contig Id
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	g136612	g2226252 g071500	g400209	g1/31129		g2619006	g126054			g1881258		g1075793		g2127359		g420808		g130130			g586808	g1945644	g225559	g2983140	:	g3257217	g129179		g2633966	Τ,	g2633967		NCBI gi
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	4.00E-124	7.30E-29	4.30E-40	3.00E-31	•	8.60E-74	7.80E-112			4.80E-64		3-131		1.10E-16		4.90E-32		3.90E-39			5.10E-35	2	4.10E-35	1.20E-33		9.70E-66	5.50E-40	•	3-276	-	2.00E-33		•
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	THYMIDYLATE SYNTHASE (TS) [Artificial gene]					(AF027868) putative L-amino acid oxidase precursor		INFLUENZAE. [Bacillus subtilis]	PRODU	(AB001	(strain MGA3) [Bacillus sp.]	aspartate kinase (EC 2.7.2.4) II precursor - Bacillus sp.	[Clostridium perfringens]	hypothetical protein 3 - Clostridium perfringens	aquaticus]		PROTEIN PHOR [Bacillus subtilis]	•	[Bacillus subtilis]	PROTEIN IN ROCR-PURA INTERGENIC REGION	HYPOTHETICAL 27.2 KD SENSORY TRANSDUCTION		ORF I	(AE000	protein	(AP000003) 597aa long hypothetical oligopeptide binding	OLIGOPEP LIDE-BINDING PROTEIN OPPA	[Bacillus subtilis]	(Z99112) chromosome segregation SMC protein homolg	[Bacillus subtilis]	_	(VEG216) [Bacillus subtilis]	
	IDYL,	4) hyp	RANE		ıs subt	7868) r	TATE	ENZA	UCTS IN SYNECHOCYSTIS AND H	1488) FUNCTION UNKNOWN, SIMILAR	MGA3	e kina	dium ₁	etical p		etical 1	IN PH	LINE	ıs subt	N N	THET!	3) hyp	231C	0692) transporter (OppBC family) [Aquifex aeolicus]	APPA [Pyrococcus horikoshii]	Ö03) 5	PEP I	is subt	2) chro	ıs subt	2) sign	16) E	•
•	ATE S	otheti	3-BOU	CAL		outativ	DEH	E E	N SY	FUNC) [Ba	se (EC	perfirin	rotein		2K pr	OR [SOHA	ilis]	ROCI	CAL:	othetic	Baci	ranspo	, F	([Bac			osomc	ilis]	al rec	acillu	
	LNA	cal pro	NDP	27.6		e L-ar	YDRO	acillus	NECE	NOIT	cillus	2.7.2	igens]	13-C		otein .	Bacill	PHAT		R-PUI	27.2 K	cal pro	llus th	orter (00000	long h			me se		ogniti	s subti	NCBI gi description
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(AF047044) putative transposase [Anabaena PCC7120] (Z99111) similar to aspartate aminotransferase [Bacillus	41 96	24 68	1.60E-11 2.40E-142	162 1392	105 1389	g3005554 g2633729	3570-3169 1146-1	1072 Bt1Gc1416 Bt1G1724 1073 Bt1Gc1414 Bt1G1725
[Bacillus cereus] (Y09476) YisO [Bacillus subtilis]	74	40	2.90E-89	891	845	g2145380	1-1293	1072 Bt1Gc1416 Bt1G1723
INTERGENIC REGION [Bacillus subtilis] (AJ000394) transcriptional regulatory protein CelR	90	98	2.00E-209	2025	2122	g4584200	1352-2647	1071 Bt1Gc1415 Bt1G1722
subtilis] HYPOTHETICAL 45.7 KD PROTEIN IN MUTT-GSIB	100 1	73	1.20E-124	1225	1603	g3025117	2875-1601	1070 Bt1Gc1412 Bt1G1721
FORMATE-LYASE 1) [Escherichia coli] (AF027868) putative alanine acetyl transferase [Bacillus	99 (53	3.80E-48	503	493	g2618995	1398-904	1070 Bt1Gc1412 Bt1G1720
[Listeria monocytogenes] FORMATE ACETYLTRANSFERASE 1 (PYRUVATE	41	68	7.60E-114	1123	1093	g129879	1979-1048	1069 Bt1Gc1413 Bt1G1719
(AJ009627) pyruvate-formate lyase activating enzyme	100 (59.	2.30E-80	807	800	g4914622	981-238	1069 Bt1Gc1413 Bt1G1718
(AB001488) ATP-DEPENDENT RNA HELICASE DEAD	75 (49	5.10E-92	917	908	g1881268	2394-3543	1068 Bt1Gc1411 Bt1G1717
(U64312) amidase [Bacillus firmus]	_	50	2.40E-117	1156	1092	g1813489	3498-2035	. 1067 Bt1Gc1410 Bt1G1716
thermoautotrophicum] (D86417) YfmI [Bacillus subtilis]	100 (33	6.10E-57	586	626	g2443252	761-1976	1067 Bt1Gc1410 Bt1G1715
subtilis] (AE000848) conserved protein [Methanobacterium	99 (32	2.30E-16	203	176	g2621778	496-56	1067 Bt1Gc1410 Bt1G1714
(AJ002571) gamma-glutamylphosphate reductase [Bacillus	100 (65	2.60E-143	1401	1401	g2632033	3166-4410	1066 Bt1Gc1409 Bt1G1713
STICKIANDIIJ GLUTAMATE 5-KINASE (GAMMA-GLUTAMYL	100 (60	7.10E-102	1010	1112	g3183548	2046-3152	1066 Bt1Gc1409 Bt1G1712
[Bacillus cereus] (AJ010739) pyrroline-5-carboxylate reductase [Clostridium	100 (56	2.40E-71	722	793	g3688282	1354-557	1066 Bt1Gc1409 Bt1G1711
(Y09212) proton /sodium-glutamate symport protein	100 (99	4.50E-155	1512	1733	g2072369	2777-1713	1065 Bt1Gc1408 Bt1G1710
(Y09212) Aspartate ammonia-lyase [Bacillus cereus]		99	5.60E-228	2200	2378	g2072370	1685-249	
INTERGENIC REGION [Bacillus subtilis] (Y09212) putative malate oxidoreductase [Bacillus cereus]	24 (92	1.60E-31	346	332	g2072371	224-1	1065 Bt1Gc1408 Bt1G1708
sp.J HYPOTHETICAL 23.7 KD PROTEIN IN ILVD-THYB	17 I	53	3.40E-08	126	122	g1730930	5367-5260	1064 Bt1Gc1406 Bt1G1707
(D90907) phosphoglycolate phosphatase [Synechocystis	100 t	34	3.60E-36	390	365	g1652645	5156-4497	1064 Bt1Gc1406 Bt1G1706
(Z95210) hypothetical protein Rv0907 [Mycobacterium	92 (24	3.60E-43	456	311	g3261758	507-3114	1064 Bt1Gc1406 Bt1G1705
NCBI gi description	Vrg	% % Ident Cvrg	BlastP- Prob	BlastP Score	nap Score	NCBI gi	Position	SEQ ID Contig Id Gene Id NO

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PHOSPHO-2-DEHYDRO-3-DEOXYHEPTONATE	100	75	9.30E-132	1292	1374	g728897	411-1482		
(L21856) repressor protein [Streptococcus pneumoniae]	- - - - - - - - - - - - - - - - - - -	6 6	7.30E-61 1.70F-130	623 1280	554 1240	g2656094 g98327	2112-3110	8 Bt1G1765	1084 Bt1Gc1428
[Bacillus subtilis] (Z94043) hypothetical protein [Bacillus subtilis]	100	58	4.70E-82	823	860	g1945713	994-1839		
(ASAL) [Methanococcus jannaschii] (Z99121) similar to maltodextrin transport system permease	79	62	2.30E-96	958	1167	g2635928	1-990	8 Bt1G1763	1084 Bt1Gc1428
	96	32	2.40E-69	703	632	g2492945	545-3017	7 Bt1G1762	1083 Bt1Gc1427
protein [Bacillus subtilis] HYPOTHETICAL 43.6 KD PROTEIN IN CPSD-METB	100	54	5.70E-109	1077	1064	g1730908	2661-1516	5 Bt1G1761	1082 Bt1Gc1425
(Z92952) product similar to Staphylococcus aureus CapA	100	31	5.20E-28	313	297	g1894741	3789-3049	4 Bt1G1760	1081 Bt1Gc1424
(Z71928) hypothetical protein [Bacillus subtilis]	100	22	2.00E-12	166	91	架495280	3818-2410	4 Bt1G1759	1081 Bt1Gc1424
(AF015609) unknown [Bacillus subtilis]	99	4	2.70E-31	344	346	g2454556	2438-1909	4 Bt1G1758	1081 Bt1Gc1424
[Archaeoglobus fulgidus]				7			-		
(AE001064) polysaccharide biosynthesis protein, putative	42	27	1.20E-12	176	95	g2650029	1728-1	4 Bt1G1757	1081 Bt1Gc1424
(AE000819) succinoglycan biosynthesis transport protein [Methanobacterium thermoautotrophicum]	. 94	23	1.20E-12	135	169	g2621399	1731-1	4 Bt1G1756	1081 Bt1Gc1424
(Z99110) yjaZ [Bacillus subtilis]	100	36	5.20E-35	379	355	g2633489	3420-2647		1080 Bt1Gc1423
(Z99110) yjbA [Bacillus subtilis]	100	70	3.20E-99	985	975	g2633495	2488-1745	3 Bt1G1754	1080 Bt1Gc1423
subtilis]			,	•	. -				
(TRYPTOPHANTRNA LIGASE) (TRPRS) [Bacillus			•						
TRYPTOPHANYL-TRNA SYNTHETASE	100	74	3.00E-135	1325	1325	g135188	469-1458	3 Bt1G1753	1080 Bt1Gc1423
(AJ0009/4) pulative S-adenosyt L-memionine:	.	40	4.90E-23	200	222	00670478	204-5/09	2C/10119 2	10/3 PHOC1422
SMALL SUBUNIT [Bacillus subtilis]		<u> </u>	200	2	3	2462060	3384 3700		
ASSIMILATORY NITRITE REDUCTASE (NAD(P)H)	99	49	5.90E-27	303	289	g1171662	3002-3319	2 Bt1G1751	1079 Bt1Gc1422
NITRITE REDUCTASE (NAD(P)H) [Bacillus subtilis]	100	57	7.50E-249	2397	2412	g1171661	581-2992	2 Bt1G1750	1079 Bt1Gc1422
[Memanococcus Jannaschii] (U57060) ScdA [Staphylococcus aureus]	53	34	6.00E-18	218	173	g1575061	1-363	2 Bt1G1749	1079 Bt1Gc1422
FERROUS IRON TRANSPORT PROTEIN B HOMOLOG	15	. 60	3.30E-29	332	332	g2498377	3584-3895	Bt1G1748	1078 Bt1Gc1421
thermoautotrophicum]		;		}	} .				
(AE000899) unknown [Methanobacterium	99	29	4.50E-06	106	8 5	g2622470	3342-3566	Bt1G1747	1078 Bt1Gc1421
IN SODA-COMGA INTERGENIC REGION PRECURSOR				٠					•
Tropa 6. acort peron	Cvrg	Ident (Prob	Score	Score	t C			
NCBI of description	%	%	BlastP-	BlastP	aat_	NCBI gi	Position	Gene Id	SEQ Contig Id

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									1431]	431		g Id
Bt1G1781 Bt1G1782 Bt1G1783	Bt1G1779 Bt1G1780	Bt1G1778	Bt1G1777	Bt1G1775 Bt1G1776		Bt1G1774	Bt1G1773	Bt1G1771	1085 Bt1Gc1431 Bt1G1770	Bt1Gc1431 Bt1G1769		Gene Id
2415-1417 3548-2640 3964-3632	818-1	4402-3464	3434-2154	5162-4578 588-16		4295-3822	2895-1148	115-321	2961-4019	1770-2877	,	Position
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g2314402 g2339996 g1841880	g2339999 g2339997	g136591	g114169	g2635179 g1168679		g3183527	g2635177	g2635175	g3123224	g399058	· .	NCBI gi
1398 752 331	348	762	1429	688 324	:	615	1224	149	869	1430	. · .	nap Score
1379 667 360	529 349	779	1461	706 309		624	1293	171	903	683		BlastP Score
5.60E-141 1.60E-65 5.40E-33	7.90	2.10	1.20E-149	1.20 1.40		5.70	7.30E-132	5.80	1.60	1.60E-115		BlastP-
.60E-141 1.60E-65 5.40E-33	6.70E-51 7.90E-32	2.10E-77	-149	1.20E-69 1.40E-27		5.70E-61	.30E-132	5.80E-13	1.60E-90	-115		•
75 47 62	3 8 51	4	66	69 38		80	43	49	.49	75		% % Ident Cvrg
100 99	99	. &	100	92: 93:	•	99	100	99	100	100		Sun %
			megaterium] 00 PUTATIVE 3-PHOSPHOSHIKIMATE 1-	subtilis] 92 (Z99117) uridine kinase [Bacillus subtilis] 93 TRANSCRIPTIONAL REPRESSOR BM3R1 [Bacillus		subtili: TRAN	10 (Z99117) similar to penicillin-binding protein [Bacillus	(Z991)	(VEGZ HISTI (IMID	CHORISMATE MUTASE [Bacillus subtilis] 00 CHORISMATE SYNTHASE (5- ENOLPYRUVYLSHIKIMATE-3-PHOSPHATE PHOSPHOLYASE) (VEGETATIVE PROTEIN 216)	ALDOLASE (PHOSPHO-2-KETO-3- DEOXYHEPTONATE ALDOLASE) (DAHP SYNTHETASE) (3-DEOXY-D-ARABINO- HEPTULOSONATE 7-PHOSPHATE SYNTHASE) /	NCBI gi description

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	1092 1	1091]	1091 1	1091 1	1091 1	·1091]		1091 1	1090 1			1090 1			1090 1))	1090 1	1090		1089 1	1089		1089 I	1088 1	SEQ NO
	Bt1Gc1432	Bt1Gc1430	Bt1Gc1430	1091 Bt1Gc1430	1091 Bt1Gc1430	1091 Bt1Gc1430		1091 Bt1Gc1430 Bt1G1793	1090 Bt1Gc1426			1090 Bt1Gc1426			1090 Bt1Gc1426		Bt1Gc1426	1090 Bt1Gc1426		1089 Bt1Gc1433	1089 BITGC1433		Bt1Gc1433	Bt1Gc1429	Contig Id
	Bt1G1799	Bt1G1798	Bt1G1797	Bt1G1796	Bt1G1795	Bt1G1794		Bt1G1793	Bt1G1792			Bt1G1791			Bt1G1790		Bt1G1789	Bt1G1788		Bt1G1787	B(101/80		Bt1G1785	Bt1G1784	Gene Id
•	1176-460	5832-5565	5550-4782	4689-4192	3836-1074	984-539		461-3	4057-3538			3439-3110			3111-2806		2722-1415	1165-80		3924-2593	19/4-1000		1-213	4334-3968	Position
	g1653574	g2633913	g1518679	g1518680	g3123287	g3183580		g2497629	g1168885			g1172708			g1172709		g732331	g2689897		g121357	g/30 044		g399378	g2339994	NCBI gi
	68	257	559	594	3685	97		539	257			260°			247	, <i>,</i>	767	636		2288	100	• 1 .	206	544	nap Score
	134	273	584	615	3712	97		377	276			280			257)	628	684		2288	1/3	j	237	470	BlastP Score
	4.90E-07	9.00E-24	9.90E-57	5.10E-60	0	4.00E-05		8.50E-35	3.40E-23			1.60E-24			4.40E-22		2.20E-61	2.50E-67		2.70E-237	3.30E-13		5.80E-20	1.20E-44	BlastP- Prob
	27	58	4	70	74	29		66	33			50			50		41	41		98	, , , , ,	3	61	8 1	% Ident
	56	99	100	99	100	99		9.	27					:	_		100	100		100	9		35	59	%Cvrg
	(D90914) negative aliphatic amidase regulator	(Z99112) similar to hypothetical proteins [Bacillus subtilis]	subtilis] (U60901) orf [Bacillus subtilis]	TRNA LIGASE) (ILERS) [Bacillus subtilis] (U60901) minicell-associated protein DivIVA [Bacillus	INTERGENIC REGION (ORF-1) [Bacillus subfilis] ISOLEUCYL-TRNA SYNTHETASE (ISOLEUCINE	HYPOTHETICAL 16.8 KD PROTEIN IN ILES-LSP	(PROLIPOPROTEIN SIGNAL PEPTIDASE) (SIGNAL PEPTIDASE/II) (SPASE II) [Bacillus subtilis]	LIPOPROTEIN SIGNAL PEPTIDASE	PUTATIVE CEL OPERON REGULATOR [Bacillus	ENZYME II, A COMPONENT) (EIII-CEL) [Bacillus subtilis]	IIA COMPONENT) (PHOSPHOTRANSFERASE	PTS SYSTEM, CELLOBIOSE-SPECIFIC IIA	ENZYME II, B COMPONENT) [Bacillus subtilis]	IIB COMPONENT) (PHOSPHOTRANSFERASE	PTS SYSTEM, CELLOBIOSE-SPECIFIC IIB	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 47.6 KD PROTEIN IN EPR-GALK	(AE000792) outer surface protein, putative [Borrelia	AMMONIA LIGASE) [Bacillus cereus]	GLUTAMINE SYNTHETASE (GLUTAMATE	305 KIBOSOMAL FROTEIN ST4 HOMOLOG [Bacillus	subtilis]	SOS REGULATORY PROTEIN LEXA/DINR [Bacillus	(Z98682) cytochrome c oxidase subunit III, aa3 type	NCBI gi description

SEQ ID Contig Id Gene Id	e Id	Position	NCBI gi	aat_ nap	BlastP Score	Table 1 BlastP- Prob	% % Ident Cvrg	Cvrg	NCBI gi description *
1092 Bt1Gc1432 Bt1G1800		2351-1497	g2982874	211	342	4.40E-31	29	100	[Synechocystis sp.] (AE000675) cobalamin synthesis related protein CobW
1092 Bt1Gc1432 Bt1G1801	1801	3196-2464	g2619006	853	. 875	1.40E-87	66	55	[Aquifex aeolicus] (AF027868) putative L-amino acid oxidase precursor
	200	166 527	~17200/2	122	140	1 405 10		B .	[Bacillus subtilis]
1093 BITOCI437 BITO1802	1802	/80-001	g1/30 943	133	148	1.500.1	20	. 4	IN UVRX-ILVA INTERGENIC REGION [Bacillus middle]
1093 Bt1Gc1437 Bt1G1803	1803	736-1911	g1001709	880	869	6.20E-87	43	100	subtilis] (D64004) hypothetical protein [Synechocystis sp.]
1093 Bt1Gc1437 Bt1G1804		3654-2830	g1170787	410	349	7.90E-32		100	OUTER MEMBRANE LIPOPROTEIN 2 PRECURSOR
1093 Bt1Gc1437 Bt1G1805	1805	4229-3676	g401471	364	213	2.00E-17	40	-60 -00	(PLP2) [Pasteurella haemolytica] HYPOTHETICAL ABC TRANSPORTER PERMEASE
1004 B+1G-1436 B+1G1806	806	1917-1148	g3557647	464	م	4 80E-48	37	3 ↔	PROTEIN YAEE [Escherichia coli]
						-			protein [Pyrococcus horikoshii]
1034 BUIGET430 BUIGIOU	1007	2/19-2931	786617CB	200	777	2.30E-18	. 0	· •	AF1627 [Archaeoglobus fulgidus]
1094 Bt1Gc1436 Bt1G1808	1808	4699-4229	g2851600	253	274	7.00E-24	35	70	HYPOTHETICAL 25.3 KD PROTEIN IN RIMI-PRFC
1095 Bt1Gc1439 Bt1G1809	1809	1-1181	g4103625	1155	1084	1.00E-109	57	87	(AF026470) gluconate permease [Pseudomonas aeruginosa]
1095 Bt1Gc1439 Bt1G1810	1810	1203-2096	g1730976	999	952	1.00E-95		100	HYPOTHETICAL 32.8 KD PROTEIN IN NUCB-AROD INTERGENIC REGION [Bacillus subtilis]
1095 Bt1Gc1439 Bt1G1811	1811	2845-2171	g732332	615	577	5.50E-56	49	100	HYPOTHETICAL 25.9 KD PROTEIN IN EPR-GALK
1095 Bt1Gc1439 Bt1G1812	1812	3099-5292	g1208461	87	113	5.90E-06	46	15	INTERGENIC REGION [Bacillus subtilis] (D64004) hypothetical protein [Synechocystis sp.]
1095 Bt1Gc1439 Bt1G1813	1813	5626-5860	g2226142	183	172	4.50E-13		45	(Y14079) hypothetical protein [Bacillus subtilis]
1096 Bt1Gc1441 Bt1G1814	1814	163-861	g399775	921	921	1.90E-92	74	100	PROBABLE MENAQUINONE BIOSYNTHESIS METHLYTRANSFERASE (SPORE GERMINATION
1096 Bt1Gc1441 Bt1G1815	1815	412-1868	g399776	1062	1056	9.50E-107	61	100	PROBABLE HEPTAPRENYL DIPHOSPHATE
				,					SYNTHASE COMPONENT II (HEPPP SYNTHASE) (SPORE GERMINATION PROTEIN C3) [Bacillus subtilis]
1096 Bt1Gc1441 Bt1G1816	1816	1993-2439	g400405	577	577	5.50E-56	72	99	NUCLEOSIDE DIPHOSPHATE KINASE (NDK) (NDP
1096 Bt1Gc1441 Bt1G1817		2706-3343	g399058	873	765	6.50E-76	77	58	CHORISMATE SYNTHASE (5-

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	Bt1Gc1446	Bt1Gc1446	Bt1Gc1446	Bt1Gc1446	Bt1Gc1447	Bt1Gc1447	Bt1Gc1447	Bt1Gc1447	Bti Gc144/	Bt1Gc1447		Bt1Gc1445	Bt1Gc1445		Bt1Gc1445	Bt1Gc1445	Bt1Gc1445	Bt1Gc1442		Bt1Gc1442			1099 Bt1Gc1442	Bt1Gc1444	} } }	Bt1Gc1443			Contig Id
	Bt1G1837	Bt1G1836	Bt1G1835	Bt1G1834	Bt1G1833	Bt1G1832	Bt1G1831	Bt1G1830	BIIGC144/ BIIG1829	Bt1G1828		Bt1G1827	Bt1G1826	•	Bt1G1825	Bt1G1824	Bt1G1823	Bt1G1822		Bt1G1821			Bt1G1820	BtlG1819		Bt1G1818		•	Gene Id
	3742-2780	2646-1618	1512-952	922-1	5074-4921	4058-4560	3563-2668	2411-2184	2019-12/9	1127-1		3535-5551	5444-3290		3487-4081	2541-3027	794-2077	3367-3049	-	2916-1780	-		179-1	4251-4545		1523-974			Position
	g2104798	g3122229	g121635	g580888	g3152725	g4584100	g2636549	g2635876	g/29450	g2635874	ंद्य	g4377189	g2635881	(g586554	g267523	g548909	g225559		g1177010	٠	. (g1730193	g2497382		g2634185			NCBI gi
	963	1123	588	1336	151	418	525	262	1082	1381		224	295		221	208	1082	501		1408			238	424		249			nap Score
•	969	1143	563	_				145	1082			275	304		309	227	_	531		1290			253	470		325			BlastP Score
	1.60E-97	5.80E-116	1.70E-54	3.50E-132	9.50E-12	2.00E-42	1.10E-43	3.30E-10	1.70E-109	2.70E-134		7.40E-24	4.60E-27		1.40E-27	6.70E-19	8.80E-104	4.10E-51	•	1.50E-131		•	1.20E-21	1.20E-44		2.80E-29			BlastP- Prob
	7 58	65	-		-	•-	-	68	80			•	7 40		7 34	32	50	97		72			78	94		38			% % Ident Cvrg
	100	100	.99	53	12	99	100	99	100	48		36	97		99	99	100	22		79	٠.		17	23	:	99		•	Cyrg %
stearothermophilus]	(Y09446) coproporphyrinogen III oxidase [Bacillus		GRPE PROTEIN [Bacillus subtilis]	(X51477) partial dnaK gene (AA 572) [Bacillus subtilis]	(AF065394) enolase [Staphylococcus aureus]	(AJ010132) hypothetical protein [Bacillus cereus]	(Z99124) similar to hypothetical proteins [Bacillus subtilis]	(Z99121) yvaL [Bacillus subtilis]	CARBOXYLESTERASE PRECURSOR [Bacillus	(Z99121) similar to hypothetical proteins [Bacillus subtilis]	[Chlamydia pneumoniae]	(AE001667) GTP Cyclohydratase & DHBP Synthase	(Z99121) similar to hypothetical proteins [Bacillus subtilis]	INTERGENIC REGION [Saccharomyces cerevisiae]	HYPOTHETICAL 21.1 KD PROTEIN IN FUS1-AGP1	HYPOTHETICAL 18.2 KD PROTEIN IN PMI 5'REGION (ORFI) (Rhizohium meliloti)	SERINE TRANSPORTER [Escherichia coli]	ORF IS231C [Bacillus thuringiensis]	INTERGENIC REGION [Bacillus subtilis]	PROBABLE RNA HELICASE IN WAPA-LICT	EPIMERASE) [Bacillus subtilis]	(GALACTOWALDENASE) (UDP-GALACTOSE 4-	UDP-GLUCOSE 4-EPIMERASE	TRANSPOSASE FOR INSERTION SEQUENCE	[Bacillus subtilis]	(Z99113) similar to thiol:disulfide interchange protein	PHOSPHOLYASE) (VEGETATIVE PROTEIN 216) (VEG216) [Bacillus subtilis]	ENOLPYRUVYLSHIKIMATE-3-PHOSPHATE	NCBI gi description

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PRECURSOR [Bacillus subtilis] transcription initiation factor sigma H - Bacillus megaterium	76 77	4.70E-66	672	639	g282367	515-1	1108 Bt1Gc1438 Bt1G1857
subtilis] IRON-UPTAKE SYSTEM BINDING PROTEIN	50 94	3.70E-73	739	749	g729483	5251-4361	1107 Bt1Gc1452 Bt1G1856
subtilis] IRON-UPTAKE SYSTEM PROTEIN FEUB [Bacillus	58 100	1.40E-91	913	1034	g729484	4331-3329	1107 Bt1Gc1452 Bt1G1855
FHUC [Bacillus subtilis] IRON-UPTAKE SYSTEM PROTEIN FEUC [Bacillus	48 100	2.10E-63	647	942	g1706787	3309-1239	1107 Bt1Gc1452 Bt1G1854
FERRICHROME TRANSPORT ATP-BINDING PROTEIN	52 100	2.10E-77	779	779	g1706796	2296-1490	1107 Bt1Gc1452 Bt1G1853
[Streptomyces antibioticus] (Z99120) yuil [Bacillus subtilis]	44 100	1.20E-60	621	556	g2635698	2398-620	1107 Bt1Gc1452 Bt1G1852
lactis] hypothetical protein 4 - Streptomyces antibioticus	47 69	1.70E-52	544	508	g419694	4630-5294	1106 Bt1Gc1450 Bt1G1851
thermoautotrophicum] (AE001272) conserved hypothetical protein [Lactococcus	35 99	1.80E-18	223	208	g3582220	3698-4039	1106 Bt1Gc1450 Bt1G1850
[Moraxella sp.] PRISMANE PROTEIN HOMOLOG [Methanobacterium	49 73	2.00E-90	902	889	g3914433	1-1051	1106 Bt1Gc1450 Bt1G1849
LIPASE 3 PRECURSOR (TRIACYLGLYCEROL LIPASE)	32 63	1.60E-14	188	135	g126303	5619-3331	1105 Bt1Gc1451 Bt1G1848
(Y14078) Hypothetical protein [Bacillus subtilis]	46 99	3.60E-11	154	162	g2226126	4317-3123	1105 Bt1Gc1451 Bt1G1847
[Clostridium perfringens] (AF027868) YoaN [Bacillus subtilis]	79 100	4.60E-178		1727	g2619026	2700-1525	1105 Bt1Gc1451 Bt1G1846
hypoth	35 <u>39</u>	2.80E-13	174	119	g2127359	1112-1	1105 Bt1Gc1451 Bt1G1845
REGULATORY PROTEIN [Bacillus subtilis]				9	9	,	
MELILOTI. [Bacillus subtilis] GI VCEROI. IIPTAKE OPERON ANTITERMINATOR	55 9 <u>8</u>	6 30E-55	567	\$56	e232174	568-1	1105 Bt1Gc1451 Bt1G1844
CATABOLISM (MOCR) GENE OF RHIZOBIUM							
(AB001488) SIMILAR TO THE RHIZOPINE	_	1.50E-163	1592	1562	g1881344	5363-3917	1104 Bt1Gc1448 Bt1G1843
PHOSPHODIESTERASE) [Bacillus subtilis] (D90905) hypothetical protein [Synechocystis sp.]	30 99	2.30E-18	222	222	g1652383	3382-3945	1104 Bt1Gc1448 Bt1G1842
PHOSPHODIESTERASE (GLYCEROPHOSPHODIESTER	_						
INTERGENIC REGION [Escherichia coli] GLYCEROPHOSPHORYL DIESTER	49 100	3.20E-74	749	721	g585208	1571-2434	1104 Bt1Gc1448 Bt1G1841
HYPOTHETICAL 50.5 KD PROTEIN IN BETT-PRPR	38 100	5.20E-83	832	763	g2495515	4251-2319	1103 Bt1Gc1449 Bt1G1840
HYPOTHETICAL 50.0 KD PROTEIN IN UNG-ROCA	33 100	9.80E-50	518	570	g732364	2170-688	1103 Bt1Gc1449 Bt1G1839
NCBI gi description	= 0			TO .	NCBI gi	Position	ID Contig Id Gene Id
	%	BlastP- %	BlastP	aat_			

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SEQ ID Contig Id Gene Id NO	Position	NCBI gi	aat_ nap Score	BlastP Score	BlastP- Prob	% % Ident Cvrg	%		NCBI gi description
1108 Bt1Gc1438 Bt1G1858	1079-570	g586909	494	502	4.80E-48	56	99	[Bacillus megaterium] HYPOTHETICAL 19.	[Bacillus megaterium] HYPOTHETICAL 19.7 KD PROTEIN IN CYSS 3'REGION
1108 Bt1Gc1438 Bt1G1859	1835-1089	g586908	940	943	9.00E-95	69	100	[Bacillus subtilis] HYPOTHETICAL TRNA/RRNA	TRNA/RRNA
								METHYLTRANS	METHYLTRANSFERASE YACO [Bacillus subtilis]
1108 Bt1Gc1438 Bt1G1860	2251-1821	g2632362	374	341	5.60E-31	55	99	(Z99104) similar t	(Z99104) similar to hypothetical proteins [Bacillus subtilis]
1108 Bt1Gc1438 Bt1G1861	2667-2236	g549024	463	378	6.70E-35	60	31	CYSTEINYL-TRI	CYSTEINYL-TRNA SYNTHETASE (CYSTEINETRNA
			.c					LIGASE) (CYSRS	LIGASE) (CYSRS) [Bacillus subtilis]
1109 Bt1Gc1454 Bt1G1862	790-1	g2632007	8 52	740	2.90E-73	63	60	(AJ002571) YkbA	(AJ002571) YkbA [Bacillus subtilis]
1109 Bt1Gc1454 Bt1G1863	3072-2284	g267022	466	513	3.30E-49	41	100	STAGE III SPOR	STAGE III SPORULATION PROTEIN J PRECURSOR
 								[Bacillus subtilis]	
1110 Bt1Gc1457 Bt1G1864	1824-1471	g2073397	553	471	9.30E-45	92	99	(Y09322) membra	22) membrane bound protein LytR [Bacillus cereus]
1110 Bt1Gc1457 Bt1G1865	3340-2165	g1709415	1306	985	3.20E-99	67	100	PYRIMIDINE NU	PYRIMIDINE NUCLEOSIDE TRANSPORT PROTEIN
1110 Bt1Gc1457 Bt1G1866	3442-3524	g225559	109	126	3.10E-07	89	6	ORF IS231C [Ba	ORF IS231C [Bacillus thuringiensis]
1111 Bt1Gc1453 Bt1G1867	338-727	g584819	184	191	4.40E-15	39	99	ATP SYNTHASE	ATP SYNTHASE PROTEIN I [Bacillus subtilis]
1111 Bt1Gc1453 Bt1G1868	735-1454	g114432	866	683	3.20E-67	70	100	ATP SYNTHASE	ATP SYNTHASE A CHAIN (PROTEIN 6) [Bacillus
1111 Bt1Gc1453 Bt1G1869	1514-1729	g114678	284	172	4.50E-13	% 1	99	megaterium] ATP SYNTHASE	megaterium] ATP SYNTHASE C CHAIN (LIPID-BINDING PROTEIN)
7.1453	1000000		}	Ì		2	3	[thermophilic bacterium PS3]	erium PS3]
1111 Bt1Gc1453 Bt1G1871	2365-2898	91168584	444 444	446	4 20E-42	46	90	ATP SYNTHASE	ATP SYNTHASE DELTA CHAIN [Bacillus megacetum]
		•	. :					stearothermophilus]	s] .
	2919-3410	g114510	773	773	9.30E-77	93	32	ATP SYNTHASE	ATP SYNTHASE ALPHA CHAIN [Bacillus megaterium]
1112 Bt1Gc1458 Bt1G1873	817-1363	g4033455	369	404	1.20E-37	45	99	SIGNAL PEPTIDASE I P (SPASE	SIGNAL PEPTIDASE I P (SPASE I) (LEADER
1113 Bt1Gc1456 Bt1G1874	1077-286	g3915949	356	343	3.40E-31	34	100	HYPOTHETICAL	HYPOTHETICAL 28.2 KD PROTEIN IN USHA-TESA
								INTERGENIC REGION []	GION []
1113 BtlGc1456 BtlG1875	2557-888	g2612897	226	319	1.20E-28	32	100	(AF015825) hypo	(AF015825) hypothetical ABC transporter [Bacillus
						}		subtilis]	;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;
1113 ptroc1430 ptro18/6	3230-220 4	71/00078	- 101	1104	3.90E-11/	0/	. 6	(299120) similar i subtilis]	(299120) Similar to NADH denydrogenase [Bacillus subtilis]
1113 Bt1Gc1456 Bt1G1877	3606-3839	g2635716	288	288	2.30E-25	68	99	(Z99120) yuzB [Bacillus subtilis]	Bacillus subtilis]
1114 Bt1Gc1460 Bt1G1878	1-1195	g2635874	1359	1335	5.20E-138	73	48	(Z99121) similar t	(Z99121) similar to hypothetical proteins [Bacillus subtilis]
1114 BtlGc1460 BtlG1879	1331-1798	g3334327	639	591	1.80E-57	76	99	SMALL PROTEI	L PROTEIN B HOMOLOG [Bacillus subtilis]
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		1125 Bt1Gc1459 Bt1G1919	1125 Bt1Gc1459 Bt1G1918	1124 Bt1Gc1469 Bt1G1917	1124 Bt1Gc1469 Bt1G1916	1124 Bt1Gc1469 Bt1G1915	1124 Bt1Gc1469 Bt1G1914		1124 Bt1Gc1469 Bt1G1913	1124 Bt1Gc1469 Bt1G1912		1123 Bt1Gc1466 Bt1G1911	. 1123 Bt1Gc1466 Bt1G1910	1122 Bt1Gc1468 Bt1G1909			1122 Bt1Gc1468 Bt1G1908		. 1122 Bt1Gc1468 Bt1G1907		1122 Bt1Gc1468 Bt1G1906		1122 Bt1Gc1468 Bt1G1905	1121 Bt1Gc1467 Bt1G1904	1121 Bt1Gc1467 Bt1G1903	1121 Bt1Gc1467 Bt1G1902	1121 Bt1Gc1467 Bt1G1901		SEQ ID Contig Id Gene Id
		2278-1	957-1	6864-6645	6599-6391	6369-5473	5447-4668		4906-3124	2792-1133		2396-1130	942-292	4073-4396			3929-2853		2491-1225		1152-616		461-6	4128-4501	3106-4119	1716-3119	228-1118		Position
		g2749982	g2894238	g2266425	g4584148	g2266426	g2266427		g2634204	g3688808		g1731090	g3170570	g225559		. •	g3123224		g1730922	.(g1730923		g1730924	g2829798	g2829797	g2829796	g2735505	,*	NCBI gi
-		87	283	178'	106	847	747		2041	1790	•	216	371	541			1126		856		661		242	201	939	1459	528		nap
·	•	136	357	194	145	850	674		2090	1835		277	395	541			1063		772		665		283	215	907	1422	565		BlastP Score
		3.20E-08	1.10E-32	1.10E-14	7.70E-09	6.40E-85	2.90E-66		2.60E-216	2.70E-189		3.40E-24	1.10E-36	3.60E-52			1.70E-107	•	1.20E-76		2.60E-65		7.80E-25	1.00E-16	5.90E-91	1.60E-145	1.00E-54		BlastP- Prob
		4	30	51	40	54	61		68	60		24	39	98		ė	59		42	٠	2		37	34	50	59	38		% % Ident Cvrg
		13	78	16	7	100	100		100	100		80	100	22			100		100		99		99	22	100	100	100		Cvrg
cDNA yk257d4.5;[]	elegans cDNA CEESX74F; coded for by C. elegans cDNA yk303f4.3; coded for by C. elegans cDNA yk257d4.3; coded for by C. elegans cDNA yk257d4.3; coded	(AF036705) Similar to phytoene desaturase; coded for by C.	(AL021841) sigJ [Mycobacterium tuberculosis]	(Y13917) yngH [Bacillus subtilis]	(AJ010111) pyruvate carboxylase [Bacillus cereus]	(Y13917) yngG [Bacillus subtilis]	(Y13917) yngF [Bacillus subtilis]	subtilis]	(Z99113) similar to propionyl-CoA carboxylase [Bacillus	(AF084104) AcsA [Bacillus firmus]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 34.6 KD PROTEIN IN GLNQ-ANSR	(AF058302) FrnE [Streptomyces roseofulvus]	ORF IS231C [Bacillus thuringiensis]	[Bacillus subtilis]	(IMIDAZOLE ACETOL-PHOSPHATE TRANSAMINASE)	HISTIDINOL-PHOSPHATE AMINOTRANSFERASE	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 48.3 KD PROTEIN IN QCRA-AROE	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 21.4 KD PROTEIN IN QCRA-AROE	INTERGENIC REGION [Bacillus subtilis]	SUBTILIS IN OCRA-AROE HYPOTHETICAL 17.3 KD PROTEIN IN OCRA-AROE	[Bacillus subtilis] TRANSPORT ATP-BINDING PROTEIN CYDC [Bacillus	[Bacillus subtilis] CYTOCHROME D UBIQUINOL OXIDASE SUBUNIT II	reductase homolog [Staphylococcus carnosus] CYTOCHROME D UBIQUINOL OXIDASE SUBUNIT I	(U96107) N5,N10-methylenetetrahydromethanopterin	nseridohiherori losis]	NCBI gi description

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TRNA LIGASE) (ASPRS) [Bacillus subtilis] HISTIDYL-TRNA SYNTHETASE (HISTIDINETRNA	68 100	1.80E-158	1544	1517	g3122900	4542-3270	· 1131 Bt1Gc1472 Bt1G1943
[Pyrococcus horikoshii] ASPARTYL-TRNA SYNTHETASE (ASPARTATE	71 100	7.10E-221	2133	2183	g3122885	3258-1486	1131 Bt1Gc1472 Bt1G1942
(AP000001) 230aa long hypothetical HESA protein	31 60	6.60E-12	161	117	g3256441	999-1	1131 Bt1Gc1472 Bt1G1941
(Z99118) similar to hypothetical proteins [Bacillus subtilis]	75 99	8.10E-62	632	622	g2635218	705-220	1131 Bt1Gc1472 Bt1G1940
sp.J PENICILLIN-BINDING PROTEINS 1A/1B (PBP1)	43 83	9.00E-150	1462	1388	g730275	5805-2078	1130 Bt1Gc1478 Bt1G1939
(D90909) quinolene resistance protein NorA [Synechocystis	28 100	1.20E-26	300	412	g1652918	3563-1525	1130 Bt1Gc1478 Bt1G1938
	48 99	1.20E-21	253	251	g1405458	1545-1261	1130 BtlGc1478 BtlG1937
acetyltransferase [Bacillus subtilis] CITRATE SYNTHASE [Bacillus coagulans]	70 76	3.70E-98	975	1017	g116465	856-1	1130 Bt1Gc1478 Bt1G1936
(Z99110) similar to ribosomal-protein-alanine N-	23 99	3.30E-10	145	74	g2633543	4794-4231	1129 Bt1Gc1474 Bt1G1935
aeruginosa] (Z93940) unknown [Bacillus subtilis]	44 99	8.50E-35	377	293	g1934836	3591-2615	1129 Bt1Gc1474 Bt1G1934
(AF147448) penicillin-binding protein 5 [Pseudomonas	31 73	4.70E-34	370	231	g4887207	1446-3762	1129 Bt1Gc1474 Bt1G1933
(AF036967) putative histidine kinase [Lactobacillus sakei]	39 76	5.70E-45	473	503	g4104606	519-1420	1129 BtlGc1474 BtlG1932
(AJ235272) unknown [Rickettsia prowazekii]	28 49	5:80E-12	182	· 8 5	g3861083	5531-2937	1128 BtlGc1475 BtlG1931
(Y13937) putative YhaQ protein [Bacillus subtilis]	39 100	3.60E-36	390	359	g2337814	2523-1861	1128 BtlGc1475 BtlG1930
(Y13937) putative YhaP protein [Bacillus subtilis]	55 1 <u>0</u> 0	9.20E-70	707	838	g2337815	1842-937	1128 Bt1Gc1475 Bt1G1929
SERINE TRANSPORTER [Escherichia coli]	46 49	5.80E-36	388	479	g548909	625-1	1128 Bt1Gc1475 Bt1G1928
LIGASE) (LEURS) [Bacillus subtilis]			1401		5012000	1020-2200	61.001.171
I FIICYI -TRNA SYNTHETASE /I FIICINETRNA			1037	1075	03173786	1625-2286	Bt1Gc1471
[Bacillus subtilis] (AF008220) YttB [Bacillus subtilis]	55 96	1.60E-96	502	1119	g2293179	1-1155	1127 Bt1Gc1471 Bt1G1926
SPOIIIAA INTERGENIC REGION [Bacillus subtilis] (Z99116) similar to hypothetical proteins from B. subtilis	67 100	9.10E-102	1009	1009	g2634887	3419-4252	1126 Bt1Gc1470 Bt1G1925
[Bacillus subtilis] HYPOTHETICAL 14.6 KD PROTEIN IN GCVT-	60 99	2.10E-38	411	403	g1731041	3242-2865	1126 Bt1Gc1470 Bt1G1924
[Aquifex aeolicus] (Z99111) similar to transcriptional regulator (LacI family)	48 100	6.30E-78	784	752	g2633758	1973-1000	1126 Bt1Gc1470 Bt1G1923
[Bacillus subtilis] (AE000776) transcriptional regulator (TetR/AcrR family)	32 48	2.40E-07	118	57	g2984362	930-1	1126 Bt1Gc1470 Bt1G1922
HYPOTHETICAL 11.9 KD PROTEIN IN HMP 3'REGION	52 73	5.30E-19	228	223	g1730269	248-1	1126 Bt1Gc1470 Bt1G1921
NCBI gi description	% at Cvrg	BlastP- % Prob Ident	BlastP Score	nap Score	NCBI gi	Position	SEQ ID Contig Id Gene Id NO

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Bt1Gc1480	1136 Bt1Gc1480 Bt1G1962	1136 BtlGc1480 BtlG1961		1135 Bt1Gc1477 Bt1G1960			1135 Bt1Gc1477 Bt1G1959		1135 Bt1Gc1477 Bt1G1958				1135 Bt1Gc1477 Bt1G1957	1134 Bt1Gc1479 Bt1G1956	1134 Bt1Gc1479 Bt1G1955		1134 Bt1Gc1479 Bt1G1954		1134 Bt1Gc1479 Bt1G1953	1133 Bt1Gc1476 Bt1G1952	1133 Bt1Gc1476 Bt1G1951	1133 Bt1Gc1476 Bt1G1950	1133 Bt1Gc1476 Bt1G1949		1133 Bt1Gc1476 Bt1G1948	1133 Bt1Gc1476 Bt1G1947		1132 Bt1Gc1473 Bt1G1946	1132 Bt1Gc1473 Bt1G1945	1132 Bt1Gc1473 Bt1G1944		NO	ID Contig Id Gene Id	SEQ
3337-2378	2425-947	1894-30		3156-2671	•		2677-1538		1526-228				136-1	5009-5751	4027-4763		2431-3801		199-2262	6026-4799	5763-4738	4103-5909	3817-2421		3503-1582	2433-276	•	5542-5841	4607-3714	3718-2480			Position	
g730786	g1304006	g1894770		g131626		(g131644		g131635				g131631	g3341855	g3341854		g732362		g124464	g421525	g586030	g1881234	g1652140		g1731309	g2635804		g420808	g2226131	g3688826		•	NCBI gi	. •
1001	1493	105		642 2	∵.		1095	•	1883	:		. •	125	564	730		1417		3426	157	1547	199	1320	;	302	75	÷	408	930	618	. - .	٠.		1. TBB
1060	1415	168		647			1124		1895		••		143	516	783		1317		3205	217	1547	222	1369		401	140		351	931	615			Score	RiastP
3.60E-107	8.60E-145	2.10E-12		2.10E-63			5.90E-114		1.20E-195				7.50E-10	1.60E-49	8.10E-78		2.10E-134		0	7.70E-18	8.90E-159	2.30E-18	6.50E-140		2.40E-37	6.10E-09		4.90E-32	1.70E-93	5.10E-60				RiastP.
61	59	30		78			57		84				61	4	60		56		93	36	83	43	54		39	35		74	59	31		1	∓	\$
	100	43		99			100		100				19	86	66		100		100	99	100	48	100		2	18		99	100	100		O	Cyrq	\$
STAGE V SPORULATION PROTEIN AE [Bacillus subtilis]	proteins and to E. coli YidA protein [Bacillus subtilis] (D84432) SpoVAF [Bacillus subtilis]	CARBOXYLASE) (AIRC) [Bacillus subtilis] (Z92954) product similar to Bacillus subtilis YxeH and YcsE	CARBOXYLASE CATALYTIC SUBUNIT (AIR	PHOSPHORIBOSYLAMINOIMIDAZOLE	CARBOXYLASE) (AIRC) [Bacillus subtilis]	CARBOXYLASE ATPASE SUBUNIT (AIR	PHOSPHORIBOSYLAMINOIMIDAZOLE	(ADENYLOSUCCINASE) (ASL) [Bacillus subtilis]	ADENYLOSUCCINATE LYASE	(VEG286A) [Bacillus subtilis]	SYNTHETASE) (VEGETATIVE PROTEIN 286A)	SUCCINOCARBOXAMIDE SYNTHASE (SAICAR	PHOSPHORIBOSYLAMINOIMIDAZOLE-	(AF077856) PotB [Actinobacillus actinomycetemcomitans]	(AF077856) PotA [Actinobacillus actinomycetemcomitans]	[Bacillus subtilis]	PROBABLE ALDEHYDE DEHYDROGENASE YWDH	thuringiensis]	IMMUNE INHIBITOR A PRECURSOR [Bacillus	orfY - Bacillus subtilis (fragment) []	SPORE PHOTOPRODUCT LYASE [Bacillus subtilis]	(AB001488) FUNCTION UNKNOWN. [Bacillus subtilis]	(D90903) glutamate decarboxylase [Synechocystis sp.]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 37.2 KD PROTEIN IN IDH-DEOR	(Z99120) gerAC [Bacillus subtilis]	aquaticus]	hypothetical 12K protein - Thermus aquaticus [Thermus	(Y14078) Hypothetical protein [Bacillus subtilis]	(AF084104) NatB [Bacillus firmus]	LIGASE) (HISRS) [Bacillus subtilis]		NCBI gi description	

Table 1

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1139 1	1139 1	1139 1	1139 1	1139 1	1138 1	1138 1	1138 1	1138 1	1138 1	1137 1	1137 1	1137 1	1137 1	1137 1	1137 1	1136 1	1136 1	NO SEQ
Bt1Gc1484	Bt1Gc1484	1139 Bt1Gc1484	Bt1Gc1484	Bt1Gc1484	Bt1Gc1485	1138 Bt1Gc1485	Bt1Gc1485	Bt1Gc1485	Bt1Gc1485	Bt1Gc1482	Bt1Gc1482	1137 Bt1Gc1482	1137 Bt1Gc1482	Bt1Gc1482	Bt1Gc1482	Bt1Gc1480	Bt1Gc1480	Contig Id
1139 Bt1Gc1484 Bt1G1981	Bt1G1980	Bt1G1979	Bt1G1978	Bt1G1977	Bt1G1976	Bt1G1975	Bt1G1974	Bt1G1973	Bt1G1972	Bt1G1971	Bt1G1970	Bt1G1969	Bt1G1968	Bt1G1967	Bt1G1966	Bt1G1965	Bt1G1964	Gene Id
4665-4360	3819-4271	2158-3600	2030-1155	120-1	6705-5951	4430-5825	4303-3881	3686-2219	1784-1251	6232-5888	5898-4831	4792-2738	2917-1413	1364-948	479-1	4646-4374	4357-3341	Position
g2636123	g1731102	g1881344	g2650107	g1731085	g1731008	g1881332	g2226205	g2352096	g1750115	g2984202	g544313	g544312	g544314	g600725	g3256494	g730784	g730785	NCBI gi
303	263	759	309	114	308	986	191	81	93	.86	542	1048	231	. 2	8 2	375	1215	nap Score
317	291	855	354	114	258	970	228	198	121	120	334	1067	307	159	159	375	1121	BlastP Score
1.90E-28	1.10E-25	1.90E-85	2.30E-32	6.30E-07	3.50E-22	1.20E-97	5.30E-19	7.90E-16	1.10E-07	4.20E-07	4.90E-52	6.50E-108	9.10E-32	1.60E-10	1.10E-11	1.40E-34	1.20E-113	BlastP- Prob
. 65	42	36	27	50	30	45	34	28	26	20	32	37	26	32	31	79	68	% Ident
99	99	100	100	25	85	100	99	100	99	4	100	100	100	19	73	60	100	Cvrg
(Z99122) ywsA [Bacillus subtilis]	MELILOTI. [Bacillus subtilis] HYPOTHETICAL 17.1 KD PROTEIN IN DNAG/DNAE 5'REGION (P17) [Listeria monocytogenes]	(AB001488) SIMILAR TO THE RHIZOPINE CATABOLISM (MOCR) GENE OF RHIZOBIUM	(AE001068) conserved hypothetical protein [Archaeoglobus	HYPOTHETICAL 17.9 KD PROTEIN IN GLNQ-ANSR INTERGENIC REGION [Bacillus subtilis]	MELILOTI. [Bacillus subtilis] HYPOTHETICAL 32.5 KD PROTEIN IN CCCA-SODA INTERGENIC REGION. [Bacillus subtilis]	(AB001488) SIMILAR TO THE RHIZOPINE CATABOLISM (MOCR) GENE OF RHIZOBIUM	(Y14082) hypothetical protein [Bacillus subtilis]	(U97022) orf; similar to serine/threonine protein	(U66480) YnaD [Bacillus subtilis]	(AE000765) flagellar biosynthetic protein Flik [Aquifex aeolicus]	[Bacillus subtilis] FLAGELLAR BIOSYNTHETIC PROTEIN FLHB [Bacillus subtilis]	[Bacillus subtilis] FLAGELLAR BIOSYNTHESIS PROTEIN FLHA	FLAGELLAR BIOSYNTHESIS PROTEIN FLHF (FLAGELLA ASSOCIATED GTP-BINDING PROTEIN)	synthetase [ryrococcus nortkosnii] (U09549) flagellar hook FlgE [Helicobacter pylori]	(AP000001) 216aa long hypothetical alanyl-tRNA	STAGE V SPORULATION PROTEIN AC [Bacillus	STAGE V SPORULATION PROTEIN AD [Bacillus	NCBI gi description

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1143	1143	1143	1143	1.142	1142	1142		1142	1142	1141		1141	1141		1141	1141	1141	1141				1140	1140			1140	1140	S E S	֓֝֝֝֟֝֝֝֓֓֓֓֓֓֓
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Bt1Gc1483	Bt1Gc1483	Bt1Gc1483	Bt1Gc1483	Bt1Gc1486	Bt1Gc1486	Bt1Gc1486		Bt1Gc1486	Bt1Gc1486	Bt1Gc1487		Bt1Gc1487	Bt1Gc1487		Bt1Gc1487	Bt1Gc148/	Bt1Gc1487	Bt1Gc1487)			Bt1Gc1481	Bt1Gc1481	`		1140 Bt1Gc1481	1140 Bt1Gc1481	Contig Id	
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Bt1G2001	Bt1G2000	Bt1G1999	Bt1G1998	Bt1G1997	Bt1G1996	Bt1G1995	-	Bt1G1994	Bt1G1993	Bt1G1992		Bt1G1991	Bt1G1990		Bt1G1989	Bt161988	Bt1G1987	Bt1G1986				Bt1G1985	Bt1G1984			Bt1G1983	Bt1G1982	Gene Id	
4813	3361	247	621	4737	3882	2196		163	6	6042		5725	5167		4853	3835	1869	139	i i			4640	2680			1770	80	Pos	
4813-3364	3361-2038	2471-1478	621-2484	4737-4253	3884-3153	2196-2008		1638-709	674-1	6042-5758		5725-5216	5167-4934		4853-3924	3839-1869	1869-1423	1395-34	i •			4640-5667	2680-4392			1770-2642	805-74	Position	
g26	g364	g215	·g177	g31:	g256	g263		g263	g263	g585949		Eg586039	g585939		g586815	g586814	g133029	g585057				g585314	g177		•	g312	g342	N	
g2649779	g3644001	g2190588	g1772644	g3150046	g2564028	g2633468		g2633470	g2633471	949		039	939		815	×14	029	057				314	g1770066			g3122275	g3425862	NCBI gi	
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453	244	251	293	371	507	168		1167	829	380		625	295		393	1973	263	1764				1121	1666			868	602	stP	
7.50	5.70	1.50	6.80	4.00	1.40	1.20		1.60E-118	1.10	4.10		4.50	4.20		1.70	6.40E-204	3.50	9.00E-182				1.20E-113	2.20E-171			8.00	1.20	BlastP Prob	
7.50E-43	5.70E-33	1.50E-20	6.80E-26	4.00E-33	1.40E-48	1.20E-12		-118	1.10E-82	4.10E-35		4.50E-61	4.20E-26		1.70E-36	204	3.50E-32	-182				-113	-171			8.00E-87	.20E-58	•	
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[Pse] (AE)	(AF)	(07:	Subs	(AF	(D&C	(Z99	Bac	(Z99	(Z99	30S	DES	SINC	30S	IN]	HYPO]	HYH	S0S	REP	RED	(AL	(AC	subtilis KETOI	(Z75	[Arc	AM	coelico PUTAT	(AJ007)		
[Pseudomonas aeruginosa] (AE001047) branched-chain amino acid ABC transporter,	[Rhodopseudomonas palustris] (AF087482) putative membrane spanning protein	(U75364) putative ABC transporter subunit	subsp. cremoris] (U62055) orfR gene product [Bacillus subtilis]	01663	(D86376) unnamed protein product [Bacillus subtilis]	(Z99109) yjzA [Bacillus subtilis]	[Bacillus subtilis]	[Dacillus subtilis] (Z99109) similar	109	RIBO	DESTABILIZING PROTEIN) [Bacillus subtilis]	SINGLE-STRAND BINDING PROTEIN (SSB) (HELIX-	30S RIBOSOMAL PROTEIN S18 (BS21) []	INTERGENIC REGION [Bacillus subtilis]	IN LENGENIC REGION [Bacillas Subills] HYPOTHETICAL 34.5 KD PROTEIN IN F	HYPOTHETICAL 74.3 KD PROTEIN IN RPLI-COTF	50S RIBOSOMAL PROTEIN L9 (BL17) []	REPLICATIVE DNA HELICASE [Bacillus subtilis]	REDUCTOISOMERASE) [Bacillus subtilis]	(ALPHA-KETO-BETA-HYDROXYLACIL	(ACETOHYDROXY-ACID ISOMEROREDUCTASE)	subtilis] KETOL-ACID REDUCTOISOMERASE	(Z75208) acetolactate synthase large subunit [Bacillus	Archaeoglobus fulgidus]	AMINOTRANSFERASE (TRANSAMINASE B) (BCAT)	coelicolor] PUTATIVE BRANCHED-CHAIN AMINO ACID	0773		•
onas a	eudon (2) pu	putat	moris orfR	<u>.</u> [] .	unna	yjzA	subtili	simik	simila	NOS		STRA	SOM	NIC	THETICAL 34.5 KD PROTEIN IN RPLI-COTF	FIIC	SOM	TIVE	OSIO	KET	TYDE	CID .	aceto)	lobus	RAN	E BR	1) 3-k		
aerugi anche	nonas tative	ive A] gene	pB ch	med p	Вас	S	er to 3	er to 3	AL P	NG P	NUE	AL P	REGI	AL 3	AL 7	AL P	DNA	MER)-BE	XXO	REDI	actate	fulgi	SFER	ANC	etoac	Z	
inosa] d-cha	palus	BC tr	produ	apero	rotei	illus s		-oxo	-oxo	ROT	ROTE	BIND	ROTI	2	4.5 K		ROT	HE	ASE)	ГА-Н	-ACI	UCTO	e synt	dus]	ASE	HED	yl-A(NCBI gi description	
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imo a	span	rter s	Bacille	molo	duct	S		acyl-	acyl-	6 (B	Baci	PRO1	18 (E	llus sı	OTE	OLE	,9 (B)	SE [cillus	ΥXO	OME	MER	large	-	ASK	Z)A rec	script	
cid A	ming	ubun	ıs sut	ğ	Baci			arrie	arrie	39) [llus s	EIN	3 S21)	ubtilis			[[7]	Bacill	subti	LAC	RORI	ASE	subur	•		M	dutase	tion	
BC t	prote	#	tilis]	actoc	llus sı		•	r prot	r prot	Bacill	übtili	(SSB		ت	RPL	- <u>-</u>		us su	lis]	2	Sudi		nit (E	i	ASE I	0 A C	nS]		
anspo	2.			occus	ubtilis			ein sy	ein sy	us su	<u>s</u>	H)(H			.00	I-CO) }	btilis]			TAS		bacille	1	B) (B	Ħ	epton		
orter,			, .	(AF016634) ClpB chaperone homolog [Lactococcus lactis	ت			[Dacinus submis] (Z99109) similar to 3-oxoacyl- acyl-carrier protein synthase	(Z99109) similar to 3-oxoacyl- acyl-carrier protein synthase	30S RIBOSOMAL PROTEIN S6 (BS9) [Bacillus subtilis]		ĽÝ.			귀	T		_			Ξ		S		CAT	5.	731) 3-ketoacyl-ACP/CoA redutase [Streptomyces		
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	1149 Bt1Gc1493 Bt1G2019 4192-3269 g2494668	1149 Bt1Gc1493 Bt1G2018 1530-1838 g1495283	1149 Bt1Gc1493 Bt1G2017 1224-2246 g4753135		1148 Bt1Gc1492 Bt1G2016 5271-4309 g730056		1148 Bt1Gc1492 Bt1G2015 3602-2265 g2792490		1148 Bt1Gc1492 Bt1G2014 2323-1950 g1731041		1148 Bt1Gc1492 Bt1G2013 1686-1495 g1731123		1148 Bt1Gc1492 Bt1G2012 1400-849 g1731125		1148 Bt1Gc1492 Bt1G2011 567-1 g1731127		1147 Bt1Gc1494 Bt1G2010 3374-4288 g3080402			1146 Bt1Gc1495 Bt1G2008 2453-2037 g2116755	·	1146 Bt1Gc1495 Bt1G2007 1-2246 g2496166	1145 Bt1Gc1489 Bt1G2006 5259-4828 g2293237	٠	1145 Bt1Gc1489 Bt1G2005 . 2535-915 g2650105			1145 Bt1Gt1489 Bt1G2004 1742-75 g2501186	1144 Bt1Gc1488 Bt1G2003 2451-735 g2145402			1144 Bt1Gc1488 Bt1G2002 796-1 g2842762	3	SEQ ID Contig Id Gene Id Position NCBI gi
	68 418	83 91	35 64		6 598		90 772		41 133		23 209		25 416		27 708		02 348		Ī	55 334		66 84	37 399		05 80			86 1854	02 1433			62 461	Score	
	449	160	168		696		730		173	•	218		445	,	717		236	•		286		243		•	164	•		1685	1456			528		BlastP Score
	2.00E-42	1.50E-11	5.30E-12		1.30E-68		3.30E-72		3.50E-13		6.00E-18		5.30E-42		8.00E-71		4.60E-38		1.10E-16	3.80E-25		1.20E-19	8.50E-35		3.20E-12			2.10E-173	3.90E-149			1.40E-49		BlastP- Prob
	_ 	42	35		45		39		33		67		4		72	•	<u>د</u>	•		43		24	54		24			63	50			4 5		% Ident
	100	35	29		100		100	•	99	•	98	٠.	99	٠	63		100		100	99		52	99		74			100	93			22	-	Cyrg %
EPIMERASE) [Methanococcus jannaschii]	PUTATIVE UDP-GLUCOSE 4-EPIMERASE	(Z71928) hypothetical protein [Bacillus subtilis]	(U52844) putative glycosyltransferase [Serratia marcescens]	PRECURSOR [Bacillus subtilis]	EXTRACELLULAR METALLOPROTEASE	[Staphylococcus aureus]	(AF041467) coenzyme A disulfide reductase	SPOIIIAA INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 14.6 KD PROTEIN IN GCVT-	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 7.0 KD PROTEIN IN BLTR-SPOIIIC	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 20.7 KD PROTEIN IN BLTR-SPOIIIC	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 32.9 KD PROTEIN IN BLTR-SPOIIIC	(Arabidoneis thalisma)	(AL022603) putative NADPH quinone oxidoreductase	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 28.2 KD PROTEIN IN GLNQ-ANSR	(D86418) YfmQ [Bacillus subtilis]	jannaschii]	HYPOTHETICAL PROTEIN MJ1236 [Methanococcus	(AF008220) YtfJ [Bacillus subtilis]	[Archaeoglobus fulgidus]		(FHS) (FTHFS) [Streptococcus mutans]	(FORMYLTETRAHYDROFOLATE SYNTHETASE)	FORMATETETRAHYDROFOLATE LIGASE	(Y09476) YitJ [Bacillus subtilis]	VITAMIN-B12 DEPENDENT) (MS) [Homo sapiens]	METHYLTRANSFERASE (METHIONINE SYNTHASE.	ATP-binding protein (braF-2) [Archaeoglobus fulgidus] 5-METHYLTETRAHYDROFOLATEHOMOCYSTEINE		NCBI gi description

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subtilis] 1153 Bt1Gc1499 Bt1G2039 5784-4865 g730774 939 971 9.70E-98 61 100 STAGE III SPORULATION PI subtilis]	1153 Bt1Gc1499 Bt1G2038 4865-4353 g1710821 500 452 9.60E-43 54 99 STAGE III SPORULATION PROTEIN AB	subtilis] 1153 Bt1Gc1499 Bt1G2037 4335-4132 g1710822 208 212 2.60E-17 56 99 STAGE III SPORULATION PROTEIN AC	1153 Bt1Gc1499 Bt1G2035 3711-2545 g1000363 1204 423 7.80E-74 59 100 (U35252) SpoIIIAE [Bacillus subtilis] 1153 Bt1Gc1499 Bt1G2036 4123-3725 g1710823 447 377 8.50E-35 66 99 STAGE III SPORULATION PROTEING	g1710825 300 272 1.10E-23 35 100	1153 Bt1Gc1499 Bt1G2032 1229-591 g1303911 335 250 2.40E-21 41 100 (D84432) SpoIIIAH [Bacillus subtilis] 1153 Bt1Gc1499 Bt1G2033 1898-1233 g1710826 426 418 3.90E-39 42 100 STAGE III SPORULATION PROTEIN AG	1152 Bt1Gc1490 Bt1G2030 3076-2492 g2275251 444 483 5.00E-46 47 100 (U44892) KdpC [Clostridium acetobutylicum] 1152 Bt1Gc1490 Bt1G2031 4450-3094 g3121784 1537 1382 2.70E-141 66 66 POTASSIUM-TRANSPORTING ATPASE B or PHOSPHOHYDROLASE [Clostridium acetob	1152 Bt1Gc1490 Bt1G2028 84-1906 g2618848 781 714 1.70E-70 40 100 (AF017113) YvmA [Bacillus subtilis] 1152 Bt1Gc1490 Bt1G2029 2725-1305 g3122316 772 871 3.80E-87 45 100 PUTATIVE SENSOR PROTEIN KDI	IN LEKGENIC REGION (ORF 1151 Bt1Gc1491 Bt1G2027 3267-3956 g2633807 738 751 2.00E-74 62 100 (Z99111) similar to ABC transp	PEPTIDASE I) [Bacillus amyloliquefaciens] 1150 Bt1Gc1496 Bt1G2025 5716-6432 g2226221 531 555 1.20E-53 46 100 (Y14082) hypothetical protein [Bacillus subtilis] 1151 Bt1Gc1491 Bt1G2026 1958-1251 g141438 355 378 6.70E-35 33 100 HYPOTHETICAL 27.3 KD PROTEIN IN TYRZ	1150 Bt1Gc1496 Bt1G2023 2521-3279 g2290993 684 697 1.10E-68 53 100 (AF006000) unknown [Bordetella pertussis] 1150 Bt1Gc1496 Bt1G2024 4981-5628 g1172045 612 639 1.50E-62 55 100 PYRROLIDONE-CARBOXYLATE PEPTII OXOPROLYL-PEPTIDASE) (PYROGLUT	1150 Bt1Gc1496 Bt1G2021 41-778 g1805478 656 659 1.10E-64 52 100 (D50453) homologue of pectin degradation of Erwinia chrysanthemi [Bacillus subtilis] 1150 Bt1Gc1496 Bt1G2022 799-2508 g1175716 944 1000 8.20E-101 39 100 HYPOTHETICAL 63.8 KD PROTEIN IN Subtilis]	ID Contig Id Gene Id Position NCBI gi nap Score Prob Ident Cvrg NO Score 1149 Bt1Gc1493 Bt1G2020 5034-4644 g2497392 668 670 7.60E-66 99 52 INSERTION SEQUENCE IS23 BINDING PROTEIN [Insertion]
subtilis] STAGE III SPORULATION PROTEIN AA [Bacillus subtilis]	ULATION PROTEIN AB [Bacillus	ULATION PROTEIN AC [Bacillus	(U35252) SpoIIIAE [Bacillus subtilis] STAGE III SPORULATION PROTEIN AD [Bacillus	SUBGINS STAGE III SPORULATION PROTEIN AF [Bacillus subtilis]	H [Bacillus subtilis] ULATION PROTEIN AG [Bacillus	P-1 (U44892) KdpC [Clostridium acetobutylicum] POTASSIUM-TRANSPORTING ATPASE B CHAIN (ATP PHOSPHOHYDROLASE [Clostridium acetobutylicum]	(Pacilius subtilis] (AF017113) YvmA [Bacillus subtilis] PUTATIVE SENSOR PROTEIN KDPD [Synechocystis	IN LEKGENIC REGION (ORF I) [Bacillus subtilis] (Z99111) similar to ABC transporter (ATP-binding protein)	PEPTIDASE I) [Bacillus amyloliquefaciens] (Y14082) hypothetical protein [Bacillus subtilis] HYPOTHETICAL 27.3 KD PROTEIN IN TYRZ-SACY	(AF006000) unknown [Bordetella pertussis] PYRROLIDONE-CARBOXYLATE PEPTIDASE (5- OXOPROLYL-PEPTIDASE) (PYROGLUTAMYL-	(D50453) homologue of pectin degradation repressor KdgR (D50453) homologue of pectin degradation repressor KdgR of Erwinia chrysanthemi [Bacillus subtilis] HYPOTHETICAL 63.8 KD PROTEIN IN SIPU-PBPC INTERGENIC BEGION [Bacillus subtilis]	NCBI gi description © INSERTION SEQUENCE IS232 PUTATIVE ATP- BINDING PROTEIN [Insertion sequence IS232]

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(Eacinus subtilis) 9 (Z99110) yjcC [Bacillus subtilis] 9 HYPOTHETICAL 7.5 KD PROTEIN IN GPSA-SPOIVA	45 99 55 99	2.90E-18 4.80E-09	221 134	195 191	g2633535 g1730916	6193-6492 6759-6959	1158 Bt1Gc1502 Bt1G2060 1158 Bt1Gc1502 Bt1G2061
- ~ ~	64 100	3.80E-119	1173	1154	g1169997	5103-6141	1158 Bt1Gc1502 Bt1G2059
	79 100	5.30E-177	1719	1838	g1730915	3774-5081	1158 Bt1Gc1502 Bt1G2058
	61 100	1.40E-82	828	927	g1730914	2574-3467	1158 Bt1Gc1502 Bt1G2057
	60 100	3.80E-64	654	654	g1730909	773-1408	1158 Bt1Gc1502 Bt1G2056
	93 30	8.40E-44	462	514	g3122814	1-346	1158 Bt1Gc1502 Bt1G2055
	44 84	9.20E-70	707	869	g3687664	4247-3080	1157 Bt1Gc1498 Bt1G2054
5'REGION (Bacillus subtilis)) 99	1.40比-41	‡	460	g11/6932	2321-3039	115/ 611061498 61102055
	_	4.40E-22	257	155	g2619048	2409-1474	Bt1Gc1498
			J.				
6 PREPROTEIN TRANSLOCASE SECA SUBUNIT	78 46	8.40E-163	1585	1585	g134396	2932-4098	1156 Bt1Gc1501 Bt1G2051
PROTEIN IN COTF-TETB INTERGENIC REGION							-
4 HYPOTHETICAL ABC TRANSPORTER ATP-BINDING	45 94	8.40E-44	462	421	g586824	1231-1894	1156 Bt1Gc1501 Bt1G2050
	85 100	4.40E-125	1229	1362	g2497856	5217-6152	1155 Bt1Gc1500 Bt1G2049
	86 100	6.00E-185	1794	1917	g2168132	3898-5172	1155 Bt1Gc1500 Bt1G2048
_	73 100	2.70E-125	1231	1413	g3123196	2628-3743	1155 Bt1Gc1500 Bt1G2047
	68 99	2.70E-38	410	522	g2293266	1746-2207	1155 Bt1Gc1500 Bt1G2046
_	43 ·100	2.20E-61	628	794	g2293327	1183-71	1155 Bt1Gc1500 Bt1G2045
INTERGENIC REGION [Bacillus subtilis]		÷					•
[Bacillus subtilis] 9 HYPOTHETICAL 137.4 KD PROTEIN IN BCSA-DEGR	29 59	9.60E-72	733	603	g1730891	5658-2476	1154 Bt1Gc1497 Bt1G2044
_	54 99	1.60E-40	431	401	g2635902	3937-2657	1154 Bt1Gc1497 Bt1G2043
INTERGENIC REGION [Bacillus subtilis]			٠		,		
coelicolor] HYPOTHETICAL 9.9 KD PROTEIN IN BCSA-DEGR	51 99	4.70E-18	219	217	g1730892	2624-2370	1154 Bt1Gc1497 Bt1G2042
	35 100	6.30E-39	416	316	g2661707	2352-1510	1154 Bt1Gc1497 Bt1G2041
8 HYPOTHETICAL P20 PROTEIN [Bacillus licheniformis]	25 98	5.90E-11	152	· 91	g141186	6270-6921	1153 Bt1Gc1499 Bt1G2040
NCBI gi description	% % Ident Cvrg	BlastP- Prob I	BlastP Score	nap Score	NCBI gi	Position	SEQ ID Contig Id Gene Id NO

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INTERGENIC REGION [Bacillus subtilis]	· <u></u>							·*. ·
	44 .45	6.00E-121	1190	1021	g1724012	6177-3207	52 Bt1Gc1507 Bt1G2078	1162
	36 100	3.70E-50	522	510	g1644225	3819-2881	52 Bt1Gc1507 Bt1G2077	1162
	63 100	2.50E-115	1137	1411	g2635701	2737-1421	52 Bt1Gc1507 Bt1G2076	1162
			•		,			
	31 100	2.80E-29	325	266	g2495406	527-2951	52 Bt1Gc1507 Bt1G2075	1162
INTERGENIC REGION [Bacillus subtilis]	* · ·							
	42 89	1.30E-22	262	233	g1176978	1-424	52 Bt1Gc1507 Bt1G2074	1162
7 (AL049485) hypothetical protein [Streptomyces coelicolor]	34 37	5.50E-08	129	68	g4539176	5646-5959	51 Bt1Gc1505 Bt1G2073	1161
S'REGION []								
	29 37	7.10E-14	186	104	g1177029	4571-2333	1161 Bt1Gc1505 Bt1G2072	110
coli]								
(TRYPTOPHANTRNA LIGASE) (TRPRS) [Escherichia								
TRYP	35 100	1.80E-41	440	348	g1711656	1344-2339	1161 Bt1Gc1505 Bt1G2071	110
[Bacillus subtilis]	·.	•					٠	
REDUCTOISOMERASE (DXP REDUCTOISOMERASE)			٠					
	58 82	2.00E-88	88 3	933	g3913544	951-1	1161 Bt1Gc1505 Bt1G2070	110
	46 100	2.10E-143	1402	1418	g1001781	3154-5287	1160 Bt1Gc1504 Bt1G2069	110
(gppA) [Helicobacter pylori 26695]				· 9.				
(AE00	26 90	2.80E-43	457	286	g2313368	1525-4004	50 Bt1Gc1504 Bt1G2068	1160
	:							
	54 99	2.70E-38	410	381	g1176526	1329-907	50 Bt1Gc1504 Bt1G2067	1160
elegans]			٠,٠.				٠.	٠.
	23 , 69	3.00E-07	138	47	g3329623	183-2148	1160 Bt1Gc1504 Bt1G2066	110
subtilis]						•		
(IMIDAZOLONEPROPIONATE HYDROLASE) [Bacillus			,.					
	72 55	7.20E-109	1076	1196	g1170427	5116-4205	1159 Bt1Gc1503 Bt1G2065	11:
PROPIONATE HYDROLASE) [Bacillus subtilis]								
	62 100	1.30E-134	1319	1334	g1170424	4198-2933	1159 Bt1Gc1503 Bt1G2064	11:
subtilis]								
(FORMIMINOGLUTAMATE HYDROLASE) [Bacillus			-					
•	49 100	7.60E-82	821	776	g1170422	2942-1980	1159 Bt1Gc1503 Bt1G2063	11:
OF MYXOCOCCUS XANTHUS. [Bacillus subtilis]			•					
_	82 99	2.10E-54	562	633	g1881322	1707-1249	1159 Bt1Gc1503 Bt1G2062	11:
INTERGENIC REGION [Bacillus subtilis]								
	Ident Cyrg	Prob 1	Score	Score	O		¢	NO E
NCBI gi description	%	ľ	BlastP	98t_	NCBI gi	Position	Contig Id Gene Id	3 E
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62 GLYCEROL-3-PHOSPHATE TRANSPORTER (G-3-P	67	1.90E-85	· 855	979	g585209	1-834	.1173 Bt1Gc1514 Bt1G2119	٠
MECE [Bacillus subtilis] 68 hypothetical protein Y - Bacillus subtilis (fragment) [Bacillus guberlic]	63	4.20E-58	597	577	g2127105	3505-4036	1172 Bt1Gc1518 Bt1G2118	_
\blacksquare	86	0	3291	3535	g586900	974-3406	1172 Bt1Gc1518 Bt1G2117	_
91 (Z82015) yukA [Bacillus subtilis]	4 8	2.80E-146	1429	1410	g1665846	5846-7543	Bt1Gc1517	_
tuberculosis]								
48 (Z95389) hypothetical protein Rv3447c [Mycobacterium	22	1.00E-41	440	266	g2104368	5644-7543	1171 Bt1Gc1517 Bt1G2115	_
100 (Z82015) yukC [Bacillus subtilis]	23	1.80E-34	374	264	g1665848	3770-5142	1171 Bt1Gc1517 Bt1G2114	_
INTERGENIC REGION [Bacillus subtilis]					·			
100 HYPOTHETICAL 36.6 KD PROTEIN IN QOXD-VPR	59	3.90E-101	1003	1015	g732353	4104-5102	1170 Bt1Gc1515 Bt1G2112	_
100 (Y11171) BC542A protein [Bacillus cereus]	96	8.50E-257	2472	2469	g2462090	3615-2161	1170 Bt1Gc1515 Bt1G2111	_
[Bacillus subtilis]			•				•	
99 (AB001488) PROBABLE ACETYLTRANSFERASE.	28	4.10E-19	229	161	g1881232	2061-1504	1170 Bt1Gc1515 Bt1G2110	
INTERGENIC REGION [Bacillus subtilis]		•						
100 HYPOTHETICAL OXIDOREDUCTASE IN PBPC-LRPC	59	2.70E-95	948	948	·g1175720	571-1470	1170 Bt1Gc1515 Bt1G2109	_
100 (AF106566) SlsA [Salmonella typhimurium]	42	2.20E-45	477	420	g4324613	2979-3656	1169 Bt1Gc1516 Bt1G2108	_
INTERGENIC REGION [Escherichia coli]							e A Paga Paga Paga Paga Paga Paga Paga Pag	
100 HYPOTHETICAL 23.1 KD PROTEIN IN DMSC-PFLA	63	2.50E-67	684	669	g146338	2997-3623	1169 Bt1Gc1516 Bt1G2107	_
					·			
100 PUTATIVE NON-HEME CHLOROPEROXIDASE	52	6.30E-71	718	721	g3914460	2503-1673	1169 Bt1Gc1516 Bt1G2106	_
INTERGENIC REGION [Bacillus subtilis]							.#	
33 HYPOTHETICAL 132.7 KD PROTEIN IN CSPB-GLPP	47	2.10E-95	949	894	g1724012	1172-1	1169 Bt1Gc1516 Bt1G2105	_
100 (Z99120) homoserine dehydrogenase [Bacillus subtilis]		6.50E-108	1067	1137	g2635723	5964-4678	1168 Bt1Gc1513 Bt1G2104	_
100 THREONINE SYNTHASE [Bacillus sp.]	9	1.10E-167	1631	1768	g135811	4679-3624	1168 Bt1Gc1513 Bt1G2103	_
pneumoniae]					,			
100 HOMOSERINE KINASE (HK) [Streptococcus	45	5.50E-63	64 3	22	g2497513	3615-2749	1168 Bt1Gc1513 Bt1G2102	_
100 LONG-CHAIN-FATTY-ACIDCOA LIGASE (LONG-		4.50E-187	1814	1805	g2829810	6115-4436	1167 Bt1Gc1509 Bt1G2101	_
99 (Z75208) hypothetical protein [Bacillus subtilis]	65	2.20E-61	628	625	g1770039	4317-3736	1167 Bt1Gc1509 Bt1G2100	_
100 (Z75208) hypothetical protein [Bacillus subtilis]	52	9.20E-54	556	648	g1770040	3715-2944	1167 Bt1Gc1509 Bt1G2099	_
subtilis]			•					
FLAVOPROTEIN SMALL SUBUNIT) (ETFSS) [Bacillus								
100 ELECTRON TRANSFER FLAVOPROTEIN BETA-	74	1.50E-83	837	977	g2494401	2862-2092	1167 Bt1Gc1509 Bt1G2098	_
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LIPOAMIDE DEHYDROGENASE COMPONENT OF		_	4.40E-189	1833	1964	g118672	4478-3069	Bt1G2143	1177 Bt1Gc1520	
PROTEIN [Bacillus subtilis]	TGL	35 100	8.00E-39	415	397	g2851451	426-1166	Bt1G2142	1177 Bt1Gc1520	
INE DECARBOXYLASE (LDC) [Bacillus subtilis]	LYS	62 23	1.10E-32	357	359	g118334	1-347	Bt1G2141	1177 Bt1Gc1520	
[Bacillus subtilis]	990)	55 33	1.90E-21	251	269	g1750108	5907-6239	Bt1G2140	1176 Bt1Gc1524	
121) similar to hypothetical proteins [Bacillus subtilis]		62 100	2.70E-54	561	674	g2635859	5739-5134	Bt1G2139	1176 Bt1Gc1524	
STAGE V SPORULATION PROTEIN K [Bacillus subtilis]		68 100	9.40E-116	1141	1119	g134777	3888-4847	Bt1G2138	1176 Bt1Gc1524	
(AF065404) pXO1-18 [Bacillus anthracis]		54 100	3.20E-90	900	897	g4894234	3780-2836	Bt1G2137	1176 Bt1Gc1524	
113) similar to host factor-1 protein [Bacillus subtilis]	(Z99	85 99	1.50E-30	337	319	g2634118	314-535	Bt1G2136	1176 Bt1Gc1524	
	[Bacillus subtilis]			-						
113) tRNA isopentenylpyrophosphate transferase	(Z99	63 30	2.90E-27	306	276	g2634117	1-271	Bt1G2135	1176 Bt1Gc1524	
INTERGENIC REGION [Bacillus subtilis]	INTERGENIC RE									
HYPOTHETICAL 73.2 KD PROTEIN IN SODA-COMGA		51 95	8.40E-163	1585	1653	g1731026	5262-3415	Bt1G2134	1175 Bt1Gc1511	
TRANSAMINASE) [Bacillus subtilis]	TRANSAMINASE									
AMINOTRANSFERASE) (D-AMINO ACID	AMINOTRANSFE									
AMINOTRANSFERASE) (D-AMINO ACID	AMINOTRANSFE									
D-ALANINE AMINOTRANSFERASE (D-ASPARTATE		45 100	7.50E-59	604	559	g3121979	2746-1893	Bt1G2133	1175 Bt1Gc1511	
						,				
18) similar to N-acetylmuramoyl-L-alanine amidase	(Z991	29 58	2.90E-29	328	228	g2635222	1679-1	Bt1G2132	1175 Bt1Gc1511	
[11] similar to hypothetical proteins [Bacillus subtilis]	(Z99)		1.40E-18		208	g2633826	7962-8166		1174 Bt1Gc1519	
[11] ykzG [Bacillus subtilis]	(Z99)		3.00E-16		202	g2633825	7414-7208			
INTERGENIC REGION [Bacillus subtilis])	}					
HYPOTHETICAL 61.5 KD PROTEIN IN ADEC-PDHA	_	87 100	8.20E-259	2491	2561	g2833392	7197-5533	Bt1G2129	1174 Bt1Gc1519	
INTERGENIC REGION (ORF4) [Bacillus subtilis]	INTERGENIC RE			٠		-				
HYPOTHETICAL 24.3 KD PROTEIN IN KINC-ADEC	_	73 100	8.50E-83	830	824	g732401	4830-5492	Bt1G2128	1174 Bt1Gc1519	
				1		0				
(AB016894) bd-type quinol oxidase subunit II [Bacillus		••	5.60E-17	151	195	g4514629	3863-5794		1174 Bt1Gc1519	
(AJ222587) YkuR protein [Bacillus subtilis]	(AJ2:		2.30E-119	1175	1244	g2632239	2495-3616	Bt1G2126	1174 Bt1Gc1519	
22587) YkuQ protein [Bacillus subtilis]	(AJ2:	80 100	6.50E-69	699	975	g2632238	1709-2416	Bt1G2125	1174 Bt1Gc1519	
22587) YkuM protein [Bacillus subtilis]	(AJ2	55 100	9.10E-86	858	823	g2632234	780-1643	Bt1G2124	1174 Bt1Gc1519	
111) ykuL [Bacillus subtilis]		56 .99	5.50E-40	426	426	g2633784	129-569	Bt1G2123	1174 Bt1Gc1519	
(AF049873) response protein [Lactococcus lactis]		55 100	2.20E-61	628	624	g3687663	4073-4738	Bt1G2122	1173 Bt1Gc1514	
04) similar to hypothetical proteins [Bacillus subtilis]	(Z99)	38 · 34	8.00E-22	260	315	g2632425	3492-1794	Bt1G2121	1173 Bt1Gc1514	
080) hypothetical protein [Bacillus subtilis]	(Y14	31 53	1.10E-09	140	121	g2226168	2836-1282	Bt1G2120	1173 Bt1Gc1514	
NSPORTER) (G-3-P PERMEASE) [Bacillus subtilis]	TRANSPORTER)									
•	·	ident Cyrg	Prob 1	Score	Score	,				
NCBI gi description		°%	1-	BlastP	nan_	NCBI gi	Position	Gene Id	ID Contig Id	
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1) similar to two-component sensor histidine kinase	(Z9911			1.20E-62	640	579	g2633724	1227-4443			
A) (SIGMA-43) [Bacillus subtilis]		8	بر مد	9 40F-29	320	530	01620924	1455-2276	Rt1G2158	1180 BriGc1526	
RNA POLYMERASE SIGMA FACTOR RPOD (SIGMA-		4 79	0 84	9.60E-130	1273	1246	g133466	5257-6135	Bt1G2157	1179 Bt1Gc1522	
5'REGION (P17) [Listeria monocytogenes] DNA PRIMASE [Bacillus subtilis]		8 100	4 48	1.80E-144	1412	1422	g130904	3389-5181	Bt1G2156	1179 Bt1Gc1522	
HYPOTHETICAL 17.1 KD PROTEIN IN DNAG/DNAE	НҮРОТ	4 99	7 ,34	5.40E-17	209	158	g1731102	2896-3336	Bt1G2155	1179 Bt1Gc1522	
DNAE INTERGENIC REGION [Bacillus subtilis]	DNAG/				i.		c				
(29) 110) similar to hypothetical proteins [bacilius subtilis] HYPOTHETICAL 30.3 KD PROTEIN IN GLYS-		100	0 71	4.10E-90	89 89 89	954	g2034937 g1731001	1716-2524	Bt1G2154	1179 Bt1Gc1522	
Scimilar to himstration proteins [Posilling subtilis]				6 20E 0	903	700	~2624057	105/ 1600	D+1C2152	1170 0+10-1522	
DNAG/DNAE INTERGENIC REGION (ORF3) [Bacillus	DNAG/I						-		-		
HYPOTHETICAL 29.3 KD PROTEIN IN BEX-		5 100	7 55	2.00E-67	685	727	g1731106	200-965	Bt1G2152	1179 Bt1Gc1522	
•					•		(
(Z99110) similar to cystathionine gamma-synthase [Bacillus		5 100	9 65	2.60E-129	1269	1248	g2633541	5178-4062	Bt1G2151	1178 Bt1Gc1521	
					į	i	q				
(Z99110) similar to cystathionine beta-lyase [Bacillus		_		1.10E-128	1263	1248	g2633542	4075-2906	Bt1G2150		
(AF031382) PagR protein [Bacillus anthracis]	•		-	5.00E-14	<u>.</u>	154	g2642588	1794-1495	Bt1G2149		
(D64000) hypothetical protein [Synechocystis PCC6803]		6 95	1 26	1.40E-34	375	240	g1001535	180-2393	Bt1G2148	1178 Bt1Gc1521	
INTERGENIC REGION [Bacillus subtilis]						!					
HYPOTHETICAL 23.2 KD PROTEIN IN SODA-COMGA		6 38	8 56	1.80E-18	223	200	g1731031	1-241	Bt1G2147	1178 Bt1Gc1521	
TOTIVE INCIDENTATION (AFGETA) [Dustings	enhtilie]						•				
TATIVE DROTEIN 220) (VEG220) [Bacillus	OFFIT	٠.									
AT BHA STIBLINIT (S COMBLEY AS ED STIBLINIT)		40	04	/.1UE-63	240	/39	83123238	/301-0830	B1102140	11// 8(1061220	
[Bacillus subtilis]			-		}	1	2	22/1 /250		1122 010 1520	
BETA SUBUNIT (S COMPLEX, 36 KD SUBUNIT)	BETA SI										
PYRUVATE DEHYDROGENASE E1 COMPONENT,		7 100	5 87	3.60E-146	1428	1477	g129068	6843-5869	Bt1G2145	1177 Bt1Gc1520	
[Bacillus subtilis]	[Bacillus	•									
COMPLEX (E2) (S COMPLEX, 48 KD SUBUNIT)	COMPL						٠			•	
COMPONENT OF PYRUVATE DEHYDROGENASE	COMPO				. ,						
DIHYDROLIPOAMIDE ACETYLTRANSFERASE		5 100	2 75	2.60E-152	1486	1564	g129054	5772-4487	Bt1G2144	1177 Bt1Gc1520	
COMPLEX, 50 KD SUBUNIT) [Bacillus subtilis]	COMPL										
(DIHYDROLIPOAMIDE DEHYDROGENASE) (S	(DIHYD										
PYRUVATE DEHYDROGENASE COMPLEX (E3)	PYRUV.							٠			
			: 	. !		Score				NO	
NCBI gi description		% % Ident Cyrg	Iden	BlastP-	BlastP Score	nap	NCBI gi	Position	Gene Id	SEQ ID Contig Id	
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(U1/283) putative spore germination apparatus protein [Bacillus megaterium]	. 100	5	6.60E-99	982	. 1190	80586018	190-1344	R1707181	1186 Bilgeissi	
[Bacillus subtilis]	99	8	2.3UE-66	0/0	6/6	g/0300/1	/699-/131			
acetobutylicum]	3 .	`	•	}			700 7171			
18 KD HEAT SHOCK PROTEIN (HSP 18) [Clostridium	. 99	37	1.20E-19	234	210	g462323	6804-6365	Bt1G2179	1185 Bt1Gc1528	
(D83967) YfkC [Bacillus subtilis]	100	42	4.30E-40	427	538	g2626815	6049-5210	Bt1G2178	1185 Bt1Gc1528	
(Y14079) hypothetical protein [Bacillus subtilis]	100	49	9.40E-84	839	888	g2226140	5004-3871	Bt1G2177	1185 Bt1Gc1528	
(Y14079) hypothetical protein [Bacillus subtilis]	100	49	2.10E-47	496	509	g2226141	3871-3230	Bt1G2176	1185 Bt1Gc1528	
5'REGION (ORFB) [Clostridium perfringens]										
HYPOTHETICAL 31.2 KD PROTEIN IN NAGH	100	32	1.20E-33	366	340	g141086	3466-2186	Bt1G2175	1185 Bt1Gc1528	
INTERGENIC REGION (ORFR) [Escherichia colii	,	. !				q	!			
HYPOTHETICAL 32.3 KD PROTEIN IN RHSE-NARV	00	29	9.70E-27	301	212	£2506666	2012-574	Bt1G2174	1185 Bt1Gc1528	
mobilis]										
ALCOHOL DEHYDROGENASE I (ADH I) [Zymomonas	%	70	1.40E-107	1064	1034	g113368	861-1	Bt1G2173	1185 Bt1Gc1528	
(D87979) YfnJ [Bacillus subtilis]	100	76	0	4166	4203	g2116974	3072-6263	Bt1G2172	1184 Bt1Gc1529	
INTERGENIC REGION [Bacillus subtilis]										
HYPOTHETICAL 58.3 KD PROTEIN IN GLPD-CSPB	79	52	1.00E-109	1084	1152	g3915990	1003-2275	Bt1G2171	1184 Bt1Gc1529	
[Bacillus subtilis]										
PROBABLE 6-PHOSPHO-BETA-GLUCOSIDASE	89	77	8.10E-165	1604	1603	g1168884	4267-3086	Bt1G2170	1183 Bt1Gc1530	
(AJ000394) ydjC-like protein [Bacillus cereus]	<u>1</u> 00	94	1.30E-104	1036	1150	g4584203	3033-2332	Bt1G2169	1183 Bt1Gc1530	
(AJ000394) chromate transport protein [Bacillus cereus]	100	93	3.10E-172	1674	1903	g4584202	1109-2287	Bt1G2168	1183 Bt1Gc1530	
(AJ000394) hypothetical protein [Bacillus cereus]	99	83	4.60E-43	455	598	g4584201	921-511	Bt1G2167	1183 Bt1Gc1530	
[Bacillus cereus]										
(AJ000394) transcriptional regulatory protein CelR	14	99	2.30E-32	354	337	g4584200	1-200	Bt1G2166	1183 Bt1Gc1530	
(AJ006589) gp35 [Bacteriophage phi-C31]	100	32	4.30E-24	276	219	g3947454	4351-3712	Bt1G2165	1182 Bt1Gc1527	
(AC006921) hypothetical protein [Arabidopsis thaliana]	61	25	1.40E-10	153	60	g4510346	7779-7200	Bt1G2164	1181 Bt1Gc1525	
(AE000757) hypothetical protein [Aquifex aeolicus]	100	23	3.50E-10	146	89	g2984109	5069-4358	Bt1G2163	1181 Bt1Gc1525	
ACETOLACTATE SYNTHASE. [Bacillus subtilis]										
(AB001488) SIMILAR TO PYRUVATE OXIDASE AND	100	54	3.30E-159	1551	1621	g1881244	4205-2478	Bt1G2162	1181 Bt1Gc1525	
FLAVOBACTERIUM SP [Bacillus subtilis]										
INDUCED PERIPLASMIC PROTEIN IN						÷.				
(AB001488) SIMILAR TO PENTACHLOROPHENOL-	9 .	8 2	1.70E-130	1280	1280	g1881356	5965-5120	Bt1G2161	1180 Bt1Gc1526	
(AF015775) YodD [Bacillus subtilis]	100	60	1.20E-65	668	654	g2415393	5089-4484	Bt1G2160	1180 Bt1Gc1526	
[Bacillus subtilis]	-	,						•		
	. (Score				NO	
NCBI gi description	(V)			Score	• 1	NCBI gi	Position	Gene Id	ID Contig Id	
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	SEQ ID Contig Id Gene Id	d Position	NCBI gi	aat_ I	BlastP Score	BlastP- Prob	% Ident C	Cvrg	NCBI gi description
	2			Score					
	1186 BtlGc1531 BtlG2182	82 3141-1777	g2828205	1579	1552	2.60E-159	66	100	PUTATIVE AMINOTRANSFERASE YHXA [Bacillus
	1186 BtlGc1531 BtlG2183	83 4875-3327	g629038	1866	1504	3.20E-154	74	100	hypothetical protein - Bacillus subtilis []
		84 5176-4871	g732347	243	119	1.90E-07	41	. 99	HYPOTHETICAL 11.7 KD PROTEIN IN EPR-GALK
					-				INTERGENIC REGION [Bacillus subtilis]
	1186 Bt1Gc1531 Bt1G2185	85 7322-4429	g3219336	81	224	1.40E-18	30	50	(AC004685) Unknown gene product [Homo sapiens]
	1186 Bt1Gc1531 Bt1G2186	86 7082-6453	g2632595	420	447	3.30E-42	40	100	(Z99105) ycgF [Bacillus subtilis]
	1186 Bt1Gc1531 Bt1G2187	87 7940-7199	g1805382	161	254	9.20E-22	26	100	(D50453) ycgG [Bacillus subtilis]
	1187 Bt1Gc1534 Bt1G2188	88 141-662	g1651966	167	190	5.60E-15	29	99	(D90901) hypothetical protein [Synechocystis sp.]
	1187 Bt1Gc1534 Bt1G2189	89 1957-3118	g1002992	316	416	6.30E-39	32	100	(U34772) ORF375 [Dichelobacter nodosus]
	1187 Bt1Gc1534 Bt1G2190	90 2698-3087	g1149666	231	229	4.10E-19	47	99	(X86498) IS1136 DNA [Clostridium perfringens]
-	1187 Bt1Gc1534 Bt1G2191	91 3239-4822	g2495662	1847	1443	9.30E-148	67 ·	95	PUTATIVE L-LACTATE PERMEASE [Escherichia coli]
	1188 Bt1Gc1532 Bt1G2192	92 554-264	g134776	454	420	2.40E-39	91	99	STAGE V SPORULATION PROTEIN G [Bacillus subtilis]
	1188 Bt1Gc1532 Bt1G2193	93 1084-710	g586881	467	469	1.50E-44	74	99	HYPOTHETICAL 13.7 KD PROTEIN IN PURR-SPOVG
	1188 Briggisso Brigo104	94 2049-1195	088880	011	045	5 50E-05	65	3 ·	DITE OPERON REPRESSOR [Racillus subtilis]
	Bt1Gc1532		g586879	1130	1001	6.40E-101	74	00	HYPOTHETICAL 31.7 KD PROTEIN IN SSPF-PURR
									INTERGENIC REGION (ORF1) [Bacillus subtilis]
	Bt1Gc1532	96 3349-3173	g1711546	296	296	3.30E-26	98	98	SSPF PROTEIN [Bacillus cereus]
	1188 Bt1Gc1532 Bt1G2197	97 3701-3444	g586231	283	284	6.10E-25	65	99	VEG PROTEIN [Bacillus subtilis]
	1188 Bt1Gc1532 Bt1G2198	98 4790-3936	g586878	902	838	1.20E-83	59	00	HYPOTHETICAL 33.3 KD PROTEIN IN KSGA-VEG
	1188 Bt1Gc1532 Bt1G2199	99 5782-4907	g585375	1044	1054	1.60E-106	70	100	DIMETHYLADENOSINE TRANSFERASE (S-
									ADENOSYLMETHIONINE-6-N', N'-ADENOSYL(RRNA) DIMETHYLTRANSFERASE) (16S RRNA
					•				DIMETHYLASE) (HIGH LEVEL KASUGAMYCIN
									RESISTANCE PROTEIN KSGA) (KASUGAMYCIN
	1188 Bt1Gc1532 Bt1G2200	00 6333-5776	e586877	659	<u>66</u>	4.20E-65	69	99	HYPOTHETICAL 20.7 KD PROTEIN IN METS-KSGA
									INTERGENIC REGION [Bacillus subtilis]
	1189 BtiGc1533 BtiG2201	01 165-1	g134763	167	119	8.20E-07	<u>ن</u>	17	SPORULATION SIGMA-E FACTOR PROCESSING PEPTIDASE (STAGE II SPORULATION PROTEIN GA)
	1189 Bt1Gc1533 Bt1G2202	02 1571-417	g120577	1421	1162	3.70E-122	78	100	[Bacillus subtilis] CELL DIVISION PROTEIN FTSZ [Bacillus subtilis]
	1189 Bt1Gc1533 Bt1G2203	03 2918-1635	g120567	1265	1223	1.90E-124	57	100	CELL DIVISION PROTEIN FTSA [Bacillus subtilis]

(Z99105) ycgF [Bacillus subtilis]	33 100		365	351	g2632595	1932-1309	1192 Bt1Gc1536 Bt1G2222	_
(AF027868) PEP synthase [Bacillus subtilis]	77 20	•	634	673	g2619033	523-1	1192 Bt1Gc1536 Bt1G2221	_
(Z79580) putative orf [Bacillus subtilis]	8 99	1.40E-18	224	301	g1620928	4602-4294	1191 Bt1Gc1538 Bt1G2220	_
NEUTRAL PROTEASE B PRECURSOR [Bacillus subtilis]	51 100	1.00E-150	1471	1386	g730171	2564-4207	1191 Bt1Gc1538 Bt1G2219	_
INTERGENIC REGION (F231) [Escherichia coli]				:	C			
HYPOTHETICAL 26.3 KD PROTEIN IN GNTR-GGT	39 100	_		3 8 7	g1176281	1706-2401		_
(AJ002571) YkoM [Bacillus subtilis]	99		116	101	g2632052	1226-1678	1191 Bt1Gc1538 Bt1G2217	_
(Z99113) similar to alkaline phosphatase [Bacillus subtilis]	59 95	7.30E-61	623	616	g2634202	1-565	1191 Bt1Gc1538 Bt1G2216	
ELEMENT IS232 [Insertion sequence IS232]								
TRANSPOSASE FOR INSERTION SEQUENCE	76 18	9.10E-26	292	286	g2497382	10032-10265	1190 Bt1Gc1535 Bt1G2215	_
(AF110737) RhsF [Sinorhizobium meliloti]	29 68	2.10E-47	496	370	g4151936	9948-8717	1190 Bt1Gc1535 Bt1G2214	_
[Archaeoglobus fulgidus]	;							
(AE000980) long-chain-fatty-acidCoA ligase (fadD-7)	28 28	2.80E-12	174	106	g2648777	10092-7477	1190 Bt1Gc1535 Bt1G2213	_
SMALL, ACID-SOLUBLE SPORE PROTEIN C5 (SASP)	71 99	3.90E-23	267	259	g134241	5213-4995	1190 Bt1Gc1535 Bt1G2212	_
•				1	(
(Z99109) similar to multidrug resistance protein [Bacillus			787	931	g2633434	4774-3504	Bt1Gc1535	_
(Y09476) YisU [Bacillus subtilis]	47 100	9.10E-47 '	490	479	g2145387	2821-3479	1190 Bt1Gc1535 Bt1G2210	
MELILOTI. [Bacillus subtilis]								
CATABOLISM (MOCR) GENE OF RHIZOBIUM			•					
(AB001488) SIMILAR TO THE RHIZOPINE	50 100	3.70E-128	1258	1233	g1881344	2742-1296	1190 Bt1Gc1535 Bt1G2209	_
[Mycobacterium leprae]					· ·.			
(Z98604) hypothetical protein MLCB2052.31	37 100	1.10E-43	461	434	g2337837	18-985	1190 Bt1Gc1535 Bt1G2208	_
subtilis]								
(X51419) internal open reading frame (AA 1-290) [Bacillus	81 100	1.70E-86	865	1251	g580938	7375-6506	1189 Bt1Gc1533 Bt1G2207	_
ACETYLOLOCOSAMINE TRANSFERASE [Bacillus								
PYROPHOSPHORYL-UNDECAPRENOL N-								
ACETYLMURAMYL-(PENTAPEPTIDE)	ř		•					
UDP-N-ACETYLGLUCOSAMINEN-	62 100	2.80E-116 (1146	1190	g585531	6399-5311	1189 Bt1Gc1533 Bt1G2206	_
DEHYDROGENASE) [Bacillus subtilis]								
REDUCTASE (UDP-N-ACETYLMURAMATE								
UDP-N-ACETYLENOLPYRUVOYLGLUCOSAMINE	66 100	3.00E-110	1089	1086	g140583	5092-4184	1189 Bt1Gc1533 Bt1G2205	_
AND SPORULATION PROTEIN) [Baçillus subtilis]	٠٠							
DIVISION INITIATION PROTEIN (CELL DIVISION	35 100	6.70E-35	378	389	g118667	4100-3303	1189 Bt1Gc1533 Bt1G2204	شر
•	it Cyrg	Prob ident	Score	Score		•	· · (7
NCBI gi description	, , ,	•	BlastP	ugn -	NCBI gi	Position	ID Contig Id Gene Id	_ 9
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protein) [Bacillus subtilis]	6	9.40E-68	9.	084	g/633300	4215-4794	1195 BttGc1537 BttGz243	_
monoc	-				81732313	317 477	Builde1337	
				ន្ទី	21730316	2105 /175	B+10-1527	_
(Z99110) oligopeptide ABC transporter (permease)	50 100	1.40E-64	658 1.	805	g2633498	2263-3195	1195 Bt1Gc1537 Bt1G2241	_
	30 10	3.40世-88	8/9	208	060C/0ìB	4/9-2128	וואס מווענוסטיי מנועעעיים	
				3	1075/07	2000	D-1627	_
COA LIGASE) (ACYL-ACTIVATING ENZYME)								
	82 63	1.20E-170	1659 1.20	1623 :	g728788	5962-4874	1194 Bt1Gc1540 Bt1G2239	
(MALTOSE TRANSACETYLASE) [Bacillus subtilis]								
	66 99		581 2.	650	g586845	4786-4232	1194 Bt1Gc1540 Bt1G2238	_
	26 99	2.10E-15	194 2.	125	g1256141	3717-3202	1194 Bt1Gc1540 Bt1G2237	_
LIGASE) (TYRRS 1) [Bacillus subtilis] .								
	71 100	3.40E-166	1617 3.40	1591	g135192	2579-1320	1194 Bt1Gc1540 Bt1G2236	_
	38 99	1.50E-12	167 1.:	178	g2633561	937-1257	1194 BtiGc1540 Bt1G2235	
	99 2	3.40E-47	494 3.4	.478	g225559	5453-5156	1193 Bt1Gc1539 Bt1G2234	_
INTERGENIC REGION [Bacillus subtilis]					•	:		
	62 99	1.50E-39	422 1.:	420	g1176959	4933-4550	1193 Bt1Gc1539 Bt1G2233	_
INTERGENIC REGION [Bacillus subtilis]							•	
0 HYPOTHETICAL 49.9 KD PROTEIN IN CITA-SSPB	57 100	4.40E-109	1078 4.40	1303	g3025180	4438-3110	1193 Bt1Gc1539 Bt1G2232	· -
				•	0	;		
				٠	g114271	2969-1545	1193 Bt1Gc1539 Bt1G2231	_
	50 92	2.00E-99	987 2.0	1226	g2495662	1-1505	1193 Bt1Gc1539 Bt1G2230	_
					(
	50 20	3.80E-31	351 3.8	328	g2633724	10228-9146	1192 Bt1Gc1536 Bt1G2229	_
	62 100	7.00E-127	1246 7.00	1284	g2108273	7744-8874	1192 Bt1Gc1536 Bt1G2228	_
[Sporosarcina halophila]			*-				• .	
	75 99	3.10E-21	249 3.	214	g134229	7478-7256	1192 Bt1Gc1536 Bt1G2227	_
	56 99	9.90E-41	433 9.9	526 .	g1945676	6970-6458	1192 Bt1Gc1536 Bt1G2226	_
0 (AF017435) orfL1 [Methylobacterium extorquens]	51 100	9.00E-63	641 9.0	662	g2394395	4897-6112	1192 Bt1Gc1536 Bt1G2225	<u>ن</u>
0 3-OXOACYL- [Thermotoga maritima]	45 100	1.20E-51	536 1.3	466	g2492749	3343-4084	1192 Bt1Gc1536 Bt1G2224	_
[Bacillus subtilis]								
9 (Z99106) transcriptional regulator (Lrp/AsnC family)	33 99	2.20E-20	241 2.3	212	g2632725	2790-3215	1192 Bt1Gc1536 Bt1G2223	_
NCBI gi description	nt Cyrg	BlastP- % Prob Ident	BlastP Bla Score Pi	aat_ Bl nap St Score	NCBI gi	Position	SEQ ID Contig Id Genė Id NO	2 - 2

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ASPARTOKINASE 1 ALPHA AND BETA SUBUNITS (ASPARTATE KINASE 1) [Bacillus subtilis]	100	67	2.70E-141	1382	1382	g416595	8279-7068	4 Bt1G2262	1198 Bt1Gc1544
DIHYDRODIPICOLINATE SYNTHASE (DHDPS) (VEGETATIVE PROTEIN 81) (VEG81) [Bacillus subtilis]	100	61	4.20E-81	. 814	926	g416876	7032-6166	4 Bt1G2261	1198 Bt1Gc1544
(Z99112) similar to hypothetical proteins [Bacillus subtilis]	100	60	3.70E-160	1560	1652	g2634050	5300-3753	4 Bt1G2260	1198 Bt1Gc1544
(M17445) ORF X [Bacillus subtilis]	100	75	2.10E-95	949	930	g143581	3635-2901	4 Bt1G2259	1198 Bt1Gc1544
STAGE III SPORULATION PROTEIN E [Bacillus subtilis]	100	ස	1.90E-225	1822	2423	g730776	2485-101	4 Bt1G2258	1198 Bt1Gc1544
SUBTILITY FENICIALIN-BINDING PROTEIN (Bacillus subtilis)	٠.					-			
STAGE V SPORULATION PROTEIN D (SPORULATION	100	69	1.80E-240	2318	2382	g586022	6406-8343	2 Bt1G2257	1197 Bt1Gc1542
subtilis]						1		•	٠
PENICILLIN-BINDING PROTEIN 2B (PBP-2B) [Bacillus	100	47	4.90E-174	1691	1623	g585648	4171-6294	2 Bt1G2256	1197 Bt1Gc1542
(U94706) cell division protein [Staphylococcus aureus]	99	27	1.70E-13	176	149	g2149892	3771-4170	2 Bt1G2255	1197 Bt1Gc1542
(ORFB) [Bacillus subtilis]						1			
HYPOTHETICAL 35.3 KD PROTEIN IN FTSL S'REGION	100	79	3.20E-122	1202	1268	g1730600	2842-3774	2 Bt1G2254	1197 Bt1Gc1542
(Z99111) yllA [Bacillus subtilis]	100	41	7.60E-114	1123	1085	g2633883	856-2469	2 Bt1G2253	1197 Bt1Gc1542
[Bacillus subtilis]									
(KETOPANTOATE REDUCTASE) (KPA REDUCTASE)				'n.					-
PROBABLE 2-DEHYDROPANTOATE 2-REDUCTASE	88	38	7.80E-41	434	386	g3913073	1-785	Bt1G2252 ,	1197 Bt1Gc1542
S'REGION []									
HYPOTHETICAL 45.4 KD PROTEIN IN THIAMINASE I	33	36	4.50E-19	233	173	g1177029	7761-7369	Bt1G2251	1196 Bt1Gc1541
[Bacillus subtilis]						• •.			
OUTB) (GENERAL STRESS PROTEIN 38) (GSP38)	÷.								-
OUTGROWTH FACTOR B) (SPORULATION PROTEIN							•		
NH(3)-DEPENDENT NAD(+) SYNTHETASE (SPORE	100	78	2.60E-111	1099	1099	g129288	6498-7313	Bt1G2250	1196 Bt1Gc1541
[Bacillus subtilis]						•			
HYPOTHETICAL 70.5 KD PROTEIN IN IDH 3'REGION	-	36	1.40E-75	762	910	g1731300	6222-4360	Bt1G2249	1196 Bt1Gc1541
(X61953) abrB [Bacillus subtilis]	99	53	2.70E-22	259	258	g39805	4226-3945	Bt1G2248	1196 Bt1Gc1541
[Bacillus sp.]									
microbial serine proteinase (EC 3.4.21) ispQ - Bacillus sp	100	58	1.20E-99	989	974	g2118109	3643-2675	Bt1G2247	1196 Bt1Gc1541
[Racillus subtilis]				•		(
(Z99106) similar to transcriptional regulator (ArsR family)	99	49	2.40E-23	269	245	£2632689	2367-2675	Bt1G2246	1196 Bt1Gc1541
(VEG147) [Bacillus subtilis]								•	
IOLS PROTEIN (VEGETATIVE PROTEIN 147)	100	45	1.30E-72	734	719	g1176985	2260-1331	Bt1Gc1541 Bt1G2245	1196 Bt1Gc154
(AB000617) YceJ [Bacillus subtilis]	<u>100</u>	49	2.80E-84	844	969	g2415747	1277-102	Bt1G2244	1196 Bt1Gc1541
TAC DI Gi description	Cyrg	Ident (Prob	Score	nap Score		r osition	Oche In	NO County to
NOBI di Apparintion	%	%	BlastP-	BlastP	88t_	NCBI di	Dosition	Cana Id	

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0 (Z99121) similar to hypothetical proteins [Bacillus subtilis] 0 (Z99121) similar to hypothetical proteins [Bacillus subtilis]	48 100 71 100		_	1762	1809	g2635916 g2635917	7292-6573 8710-7280	Bt1Gc1546 Bt1G2278 Bt1Gc1546 Bt1G2279	1201 B
Bacillu					,				
PEPTID			9.70		Ų	6110700	0,00		
INTERGENIC REGION [Bacillus subtilis]	35 100	8 40F-28 3	x 40	<u></u>	356 .	g116906	6460-5717	Rt1Gc1546 Rt1G2277	1201 P a
	48 100	8.50E-51 4	8.50	528	24	g732355	4538-5305	Bt1Gc1546 Bt1G2276	1201 B
	73 96	_	٠		956	g4.160468	3541-5356		
						•	-		
[Bacillus megaterium] 0 (Z99119) similar to Na+-transporting ATP synthase	53 100	6.50E-92 5		916	1192	g2635594	1911-3242	Bt1Gc1546 Bt1G2274	1201 Bi
8 SMALL, ACID-SOLUBLE SPORE PROTEIN C3 (SASP)	72 98	4.60E-20 7	4.60	238	229	g134237	1626-1820	Bt1Gc1546 Bt1G2273	1201 B
9 (D88825) ORF3 [Aeromonas caviae]	1 99	9.00E-24 41	9.00	273	267	g2335053	1071-1472	Bt1Gc1546 Bt1G2272	1201 B
(AF109	4 80	2.00E-35 64	2.00	383	399	g4160472	1-354	Bt1Gc1546 Bt1G2271	1201 B
licheniformis]									
(AF007	5 43	0 . 35		4102	3841	g2982196	1-8387	Bt1Gc1543 Bt1G2270	1200 B
NUSB			9		į	6			
	54 99	6 70F_35 S	6 70	378	343	01709418	6346-5948	Rt1Gc1545 Rt1G2269	1199 B
METHENYLTETRAHYDROFOLATE								•	
DEHYDROGENASE /					•	•			
	7 100	3.70E-98 67	3.70	975	1003	g1706886	5930-5082	1199 Bt1Gc1545 Bt1G2268	1199 B
[Bacillus subtilis]									
	54 100		1.30E-118	1168	1226	g1706723	5046-3700	1199 Bt1Gc1545 Bt1G2267	1199 BI
									: }
SUBUNIT (EXONUCLEASE VII SMALL SUBUNIT)					٠,				. •
	4 98	3.60E-11 54	3.60	154	147	g1706724	3620-3459	Bt1Gc1545 Bt1G2266	11 99 Bı
[Bacillus stearothermophilus]				٠٠.					
DIPHOSPHATE SYNTHASE) (FPP SYNTHASE)									
	7 100	1.40E-80 57	1.40	809	865	g585326	3468-2578	Bt1Gc1545 Bt1G2265	1199 B1
SYNTHASE (DXP SYNTHASE) [Bacillus subtilis]						!			
	7 100)-267 77	5.70E-267	2568	2603	g1731052	2273-376	Bt1Gc1545 Bt1G2264	1199 Bt
INTERGENIC REGION [Bacillus subtilis]		1.000			407	81/21101	3//-1	מניסביטיום פרנסביסט	
				Æ 474	763	~1731101	377 1		9
NCBI gi description	t Cyrg	P- % b Ident	Prob	Score	nap Score	NCBI gi	Position	Contig Id Gene Id	
					88t				SEO

		٠			Table 1			
SEQ ID Contig Id Gene Id NO	Position	NCBI gi	nap Score	BlastP Score	BlastP- Prob	% Ident	% Cvrg	NCBI gi description
1201 Bt1Gc1546 Bt1G2280	9446-8733	g1945722	950	950	1.60E-95	74	100	(Z94043) hypothetical protein [Bacillus subtilis]
1201 Bt1Gc1546 Bt1G2281	9973-9615	g2635931	230	215	1.30E-17	43	68	(Z99121) similar to transcriptional regulator (GntR family) [Bacillus subtilis]
1202 Bt1Gc1547 Bt1G2282	3544-1338	g2493678	97	218	7.70E-17	25	. 35	OVIDUCT-SPECIFIC GLYCOPROTEIN PRECURSOR (OVIDUCTAL GLYCOPROTEIN) (OVIDUCTIN)
							41.4	musculus]
1202 Bt1Gc1547 Bt1G2283	4729-1097	g2444121	& 3	192	7.70E-14	36	10	(U88974) ORF42 [Streptococcus thermophilus temperate
1202 B+1C+1522 B+1C2204	7 1427	2700077	1330	1364	7 EVE 130	'n	3	bacteriophage O1205]
		9	i	Ċ	1000	Ş		PHOSPHATE DEHYDROGENASE (NON-
	•			•				PHOSPHORYLATING GLYCERALDEHYDE 3-
	-	٠					•	PHOSPHATE DEHYDROGENASE)
,	•							(GLYCERALDEHYDE-3-PHOSPHATE
								DEHYDROGENASE) [Streptococcus mutans]
1203 Bt1Gc1523 Bt1G2285	4524-2737	g1709184	91	186	1.50E-14	37	40	GLUTAMATE RACEMASE [Bacillus sphaericus]
1203 Bt1Gc1523 Bt1G2286	3391-4797	g2635847	1477	1483	5.40E-152	59	100	(Z99121) similar to ABC transporter (amino acid permease) [Bacillus subtilis]
1203 Bt1Gc1523 Bt1G2287	4876-6074	g1706578	469	448	2.60E-42	29	100	PROTEIN ECSB [Bacillus subtilis]
1203 Bt1Gc1523 Bt1G2288	6248-6453	g231698	153	195	9.10E-15	59	15	CATALASE []
Bt1Gc1550	327-1	g225559	503	515	2.00E-49	94	23	ORF IS231C [Bacillus thuringiensis]
Bt1Gc1550	717-4011	g128494	67	148	3.30E-09	27	12	NODULATION PROTEIN V [Bradyrhizobium japonicum]
Bt1Gc1550	1-2399	g2633696	816	934	8.10E-94	35	76	(Z99110) similar to hypothetical proteins [Bacillus subtilis]
Bt1Gc1550	1062-3089	g1763702	53	147	1.00E-08	24	69	(Z83337) similar to phosphatases [Bacillus subtilis]
1205 Bt1Gc1549 Bt1G2293	2777-3964	g585035	1214	1145	3.50E-116	62	. 100	PENICILLIN-BINDING PROTEIN DACF PRECURSOR (D-ALANYL-D-ALANINE CARBOXYPEPTIDASE) (DD-
		:		• • •				PEPTIDASE) (DD-CARBOXYPEPTIDASE) [Bacillus subcilici
1205 Bt1Gc1549 Bt1G2294	4137-4484	g3287912	431	431	1.60E-40	72	99	ANTI-SIGMA F FACTOR ANTAGONIST (STAGE II
1205 B+1C-15/10 B+1C2205	\$ COV 88VV	~12/757	5 01	6 01	1 005 47	70	9 :	SPORULATION PROTEIN AA) [Bacillus coagulans] ANTI SIGMA E FACTOR (STACE II SECRITI ATTOM
ع		C	:	;		;	- 1	PROTEIN AB) [Bacillus licheniformis]
1205 Bt1Gc1549 Bt1G2296	4941-5699	g464690	1071	. 1023	3.00E-103	83	100	RNA POLYMERASE SIGMA-F FACTOR (STAGE II SPORTII ATTON PROTEIN AC) (SPORTII ATTON SIGMA
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 1210 Bt1Gc1555 Bt1G2320	1210 Bt1Gc1555 Bt1G2319	1210 Bt1Gc1555 Bt1G2318	1210 Bt1Gc1555 Bt1G2317	1210 Bt1Gc1555 Bt1G2316	1209 Bt1Gc1556 Bt1G2315		1209 Bt1Gc1556 Bt1G2314	1208 Bt1Gc1552 Bt1G2313	1208 Bt1Gc1552 Bt1G2312	ā	1208 Bt1Gc1552 Bt1G2311		1208 Bt1Gc1552 Bt1G2310	1208 Bt1Gc1552 Bt1G2309		1208 Bt1Gc1552 Bt1G2308	1208 Bt1Gc1552 Bt1G2307	1208 Bt1Gc1552 Bt1G2306	1208 Bt1Gc1552 Bt1G2305	1207 Bt1Gc1551 Bt1G2304		1207 Bt1Gc1551 Bt1G2303		1207 Bt1Gc1551 Bt1G2302	1206 Bt1Gc1548 Bt1G2301	1206 Bt1Gc1548 Bt1G2300	1206 Bt1Gc1548 Bt1G2299	1205 Bt1Gc1549 Bt1G2298	1205 Bt1Gc1549 Bt1G2297	Č	SEQ ID Contig Id Gene Id
5129-5434	5016-4415	3669-4409	3526-2762	218-1021	3833-5640		639-3659	9596-9487	9233-7520		7462-8475		8931-6873	5782-4124		4061-3630	3069-2433	1702-617	400-1	4121-4587		1586-1992		320-1	4331-3361	3067-2210	1965-554	7572-8172	5832-7304	٠.	Position
g2635728	g2635729	g1592700	g2226236	g2635730	g3123076		g2622469	g225559	g3043927		g1731075		g2935421	g468461		g2634168	g1001645	g1929335	g2612891	g2370586		g3257497		g2497382	g114510	g114636	g728922	g730782	g2648543		NCBI gi
377	189	478	422	1155	297		1087	155	212		1158		294	761	-	444	201	1142	306	85		63		532	1429	1052	2035	419	1132	2016	
390	242	528	416	1162	276	•	1282	161	259		1162		355	908		462	213	.1149	311	133		126	z+	548	1413	1070	1969	298	1006		BlastP Score
3.60E-36	1.70E-20	8.50E-51	6.30E-39	5.60E-118	4.30E-24		1.10E-130	5.10E-11	2.70E-22		5.60E-118		1.80E-32	4.60E-91		8.40E-44	2.00E-17	1.30E-116	8.40E-28	6.10E-09		3.40E-08		6.50E-53	1.40E-144	3.10E-108	1.70E-203	2.00E-26	1.90E-101		BlastP- Prob
62	27	40	38	82	34		43	86	34	٠	65		49	36	••.	59	28	56	41	4		27		99	%	72	86	4 3	47		% Ident
99	100	100	100	100	87		97	00	85		100	٠٠,	38	100		99	100	100	78			99		25	2	100	100	100	100		Cvrg
(Z99120) yutD [Bacillus subtilis]	(Z99120) yutC [Bacillus subtilis]	(Y08559) Unknown [Bacillus subtilis]	(Y14083) hypothetical protein [Bacillus subtilis]	[Synechocystis sp.] (Z99120) similar to lipoic acid synthetase [Bacillus subtilis]	HYPOTHETICAL 28.9 KD PROTEIN SLL0617	[Methanobacterium thermoautotrophicum]	(AE000899) ferrous iron transport protein B	ORF IS231C [Bacillus thuringiensis]	(AF039103) Tat-interacting protein TIP30 [Homo sapiens]	OXIDOREDUCTASE YQJM [Bacillus subtilis]	PROBABLE NADH-DEPENDENT FLAVIN	[Pseudoalteromonas sp. S9]	(AF047839) adaptive response regulatory protein	(D28859) TraC [Enterococcus faecalis]	subtilis]	(Z99113) similar to fosfomycin resistance protein [Bacillus	(D64002) hypothetical protein [Synechocystis sp.]	(Z93767) ywrH [Bacillus subtilis]	(AF015825) MutT homolog [Bacillus subtilis]	(Y11313) hypothetical protein [Xanthomonas campestris]	horikoshii]	(AP000004) 132aa long hypothetical protein [Pyrococcus	ELEMENT IS232 [Insertion sequence IS232]	TRANSPOSASE FOR INSERTION SEQUENCE	ATP SYNTHASE ALPHA CHAIN [Bacillus megaterium]	ATP SYNTHASE GAMMA CHAIN [Bacillus megaterium]	subtilis] ATP SYNTHASE BETA CHAIN [Bacillus caldotenax]	[Archaeoglobus fulgidus] STAGE V SPORULATION PROTEIN AA [Bacillus	(AE000965) sodium- and chloride-dependent transporter		NCBI gi description

	1213 Bt1Gc1557 Bt1G2340 8715-6985	1213 Bt1Gc1557 Bt1G2339 3122-1896	1213 Bt1Gc1557 Bt1G2338 1903-1346	1213 Bt1Gc1557 Bt1G2337 834-391	1212 Bt1Gc1553 Bt1G2336 7963-7369	1212 Bt1Gc1553 Bt1G2335 7256-6585	1212 Bt1Gc1553 Bt1G2334 6213-4875	1212 Bt1Gc1553 Bt1G2333 4735-3194	1212 Bt1Gc1553 Bt1G2332 3064-2011		1212 Bt1Gc1553 Bt1G2331 1771-2016	, a	1212 Bt1Gc1553 Bt1G2330 1593-757	1211 Bt1Gc1554 Bt1G2329 6133-6/80		1211 Bt1Gc1554 Bt1G2328 5337-6148	1211 Bt1Gc1554 Bt1G2327 5800-4288		1211 Bt1Gc1554 Bt1G2326 3808-4258	1211 Bt1Gc1554 Bt1G2325 2951-3709	1211 Bt1Gc1554 Bt1G2324 1165-2922	 1211 Bt1Gc1554 Bt1G2323 525-1130	1211 Bt1Gc1554 Bt1G2322 230-1	1210 Bt1Gc1555 Bt1G2321 6176-4268	Contig Id	SEO
	5 g2829799	_		g586818	9 g1730976			-		ļ	5 g1176942	(g586827) g12/250			-		8 g140720	g118613	2 g1071812	g118620	g1176915	\$ g2501678	NCBI gi	
	1397	397: ₅	407	291	842	511	1017	1712	1106		137		819	526	}	692	508		353	1160	2563	81 1.	168	. 71	nap Score	99†
	1397 - 1379		421	296	844	547	754	1706	1142		154		775	44		668	570	· •	393	1169	2570	819	200	130	Score	! i
	5.60E-141	4.80E-48	1.90E-39	3.30E-26	2.80E-84	8.30E-53	9.60E-75	1.30E-175	7.30E-116		3.60E-11		5.70E-77	5.30E-42		1.20E-65	3.00E-55		. 1.70E-36	1.00E-118	3.50E-267	1.20E-81	4.90E-16	4.80E-08	Prob	! i
	47	30	4	5 37	82	51	46	2	62		39		55	.		51	4		50	82	82	71	4	27	% Ident	;
	100	100	.99	99	67	100	100	100	100		99		100	28		100	100		99	100	100	100	52	30	Cvrg	?
subtilis]	[Mycobacterium tuberculosis] TRANSPORT ATP-BINDING PROTEIN CYDD [Bacillus		INTERGENIC REGION [Bacillus subtilis] rfbO protein - Vibrio cholerae [Vibrio cholerae]	INTERGENIC REGION [Bacillus subtilis] HYPOTHETICAL 16.0 KD PROTEIN IN COTF-TETB	HYPOTHETICAL 32.8 KD PROTEIN IN NUCB-AROD		subtilis] (AF026470) gluconate permease [Pseudomonas aeruginosa]	GLUCONOKINASE (GLUCONATE KINASE) [Bacillus	(AJ002	INTERGENIC REGION (ORF3) [Rhizobium meliloti]	HYPOTHETICAL 8.9 KD PROTEIN IN UREA-UREB	INTER		CHEMOTAXIS MOTB PROTEIN (MOTILITY PROTEIN B) [Bacillus subtilis]	A) [Bacillus subtilis]	CHEMOTAXIS MOTA PROTEIN (MOTILITY PROTEIN	(AF006665) YokZ [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis]	PROTEIN [Bacillus subtilis] HYPOTHETICAL 17.1 KD PROTEIN IN SDHB-GERE	Bacillus subtilis [] SUCCINATE DEHYDROGENASE IRON-SULFUR	succinate dehydrogenase (EC 1.3.99.1) flavoprotein -	"REGION [Bacillus subtilis] SUCCINATE DEHYDROGENASE CYTOCHROME B-	HYPOTHETICAL 17.3 KD PROTEIN IN LYSC	HYPOTHETICAL 33.7 KD PROTEIN IN LSP-PYRR INTERGENIC REGION (ORF-X) []	NCBI gi description	

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FHUB [Bacillus subtilis]	-							•
FERRICHROME TRANSPORT PERMEASE PROTEIN	33 89	2.60E-24	278	362	g1706795	1-1476	Bt1G2359	1217 Bt1Gc1559
INTERGENIC REGION [Bacillus subtilis]					. (٠	
HYPOTHETICAL 79.2 KD PROTEIN IN PHOH-DGKA	46 87	2.00E-138		1488	g1730999	4439-6303	Bt1G2358	1216 Bt1Gc1564
(AJ224829) ORF4 [Bacillus megaterium]	73 100	1.10E-121	1197	1194	g3947511	3297-4268	Bt1G2357	1216 Bt1Gc1564
(AJ224829) sporulation protein [Bacillus megaterium]	50 100	3.80E-96	956	1071	g3947510	2094-3278	Bt1G2356	1216 Bt1Gc1564
(AJ224829) ORF3 [Bacillus megaterium]		•	274	333	g3947509	1633-1914	Bt1G2355	
INTEREGENIC REGION [Bacillus subtilis]								
HYPOTHETICAL 16.8 KD PROTEIN IN RPSU-PHOH	67 99	9.00E-24	273	477	g1730992	276-716	Bt1G2354	1216 Bt1Gc1564
subtilis]			,					
30S RIBOSOMAL PROTEIN S21 (BS-B) [Bacillus	98 98	1.80E-16	204	272	g1710750	87-257	Bt1G2353	1216 Bt1Gc1564
MELILOTI. [Bacillus subtilis]								
CATABOLISM (MOCR) GENE OF RHIZOBIUM								
(AB001488) SIMILAR TO THE RHIZOPINE	36 81	1.40E-64	658	593	g1881344	7684-6516	Bt1G2352	1215 Bt1Gc1561
(Z99124) similar to hypothetical proteins [Bacillus subtilis]	36 100	9.10E-32	217	461	g2636549	6485-5596	Bt1G2351	1215 Bt1Gc1561
(D86418) YfmQ [Bacillus subtilis]	47 99	9.80E-34	367	339	g2116755	5410-4992	Bt1G2350	1215 Bt1Gc1561
subtilis (fragment) [Bacillus subtilis]								
hypothetical protein (212 aa) (ponA operon) - Bacillus	44 100	9.90E-41	433	444	g2126959	3531-4184	Bt1G2349	1215 Bt1Gc1561
Methanococcus jannaschii [Methanococcus jannaschii]								-
capsular polysaccharide biosynthesis protein D homolog -	43 100	1.70E-68	695	620	g2127789	3130-2149	Bt1G2348	1215 Bt1Gc1561
aeruginosa]			•					
(AF009955) oxidoreductase Rmd [Pseudomonas	29 89	1.10E-23	272	174	g3249547	1927-1222	Bt1G2347	1215 Bt1Gc1561
pneumoniae]			٠					
UDPGIcNAc GlcNAc transferase [Streptococcus	ن							. •
(AF068902) undecaprenyl-PP-MurNAc-pentapeptide-	38 7.7	1.10E-48	508	451	g4009481	815-1	Bt1G2346	1215 Bt1Gc1561
ELEMENT IS232 [Insertion sequence IS232]								
TRANSPOSASE FOR INSERTION SEQUENCE	99 24	1.70E-52	544	528	g2497382	6453-6769	Bt1G2345	1214 Bt1Gc1558
MONOOXYGENASE) []								•
PHENYLALANINE-4-HYDROXYLASE (PAH) (PHE-4-	29 100	1.10E-36	395	299	g1172469	3412-2433	Bt1G2344	.1214 Bt1Gc1558
3'REGION [Bacillus subtilis]		•						
HYPOTHETICAL 31.5 KD PROTEIN IN GLVBC	42 100	2.10E-54	562	475	g1723610	446-1285	.Bt1G2343	1214 Bt1Gc1558 Bt1G2343
[Bacillus subtilis]		-					٠	
HYPOTHETICAL PROTEIN IN GLVBC 3'REGION	38 99	4.10E-12	163	199	g1723608	11-388	Bt1G2342	1214 Bt1Gc1558
	01	7.11-		1101	84047170	10000-0700	14670110	יייייייייייייייייייייייייייייייייייייי
TO ANICOORT ATO DIMININIC DECITION CVING COSSILIA	70		2	1104	~7°7070°	10060 8706	B+1/C22/1	IJ
NCBI gi description	\circ	ī		nap Score	NCBI gi	Position	Gene Id	NO Config to
	%	BlastP- %	BlastP		NCDI :	J) L	

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		0.078	87	231	g1076839	7425-5506	Bt1Gc1565	
(UDP-N-ACETYLMURANOYL-L-ALANINE SYNTHETASE) [Bacillus subtilis] (AF008220) YtoT [Bacillus subtilis]	54 100	1.60E-184	1790	1750	£2293215	6687-4105	1221 Bt1Gc1565 Bt1G2380	
	74 100	8.00E-174	1689	1685	g3183540	2654-1359	1221 Bt1Gc1565 Bt1G2379	
(AB001488) PROBABLE ACETYLTRANSFERASE.	40 99	2.40E-30	335	364	g1881232	1161-616	1221 Bt1Gc1565 Bt1G2378	
	44 55	3.70E-82	824	782	g2226165	7346-6222	1220 Bt1Gc1560 Bt1G2377	
	55 100	3.30E-49	513	715	g1945085	2821-1055	1220 Bt1Gc1560 Bt1G2376	
INTERGENIC REGION [Bacillus subtilis]			•		,			
HYPOTHETICAL 25.8 KD PROTEIN IN EPR-GALK	50 100	3.30E-42	447	559	g732337	246-964	1220 Bt1Gc1560 Bt1G2375	
(AF008220) putative malolactic enzyme [Bacillus subtilis]	70 90	3.70E-169	1645	1859	g2293209	7297-5768	1219 Bt1Gc1563 Bt1G2374	
	44 100	5.00E-62	634	824	g3599667	5579-4447	1219 Bt1Gc1563 Bt1G2373	
	91 100	8.00E-142	1387	1440	g1769946	4384-3455	1219 Bt1Gc1563 Bt1G2372	
	93 100	0	5081	5149	g1769947	3333-142	1219 Bt1Gc1563 Bt1G2371	
(AF008220) YtcG [Bacillus subtilis]	75 99	2.80E-52	542	619	g2293279	7460-7005	1218 Bt1Gc1562 Bt1G2370	
ATTACHMENT PROTEIN [Bacillus subtilis]				•			,	
REPLICATION INITIATION AND MEMBRANE	47 100	9.70E-114	1122	1038	g118711	6843-5452	1218 Bt1Gc1562 Bt1G2369	
	51 100	2.60E-81	816	806	g140025	5403-4468	1218 Bt1Gc1562 Bt1G2368	
INTERGENIC REGION (ORF-281) [Bacillus subtilis]								
HYPOTHETICAL 33.3 KD PROTEIN IN DNAI-THRS	31 100	1.00E-38	414	378	g140017	3998-3141	.1218 Bt1Gc1562 Bt1G2367	
TRNA LIGASE) (THRRS) [Bacillus subtilis]								
THREONYL-TRNA SYNTHETASE I (THREONINE	80 100	6.20E-277	2662	2706	g135175	2809-876	1218 Bt1Gc1562 Bt1G2366	
(Z75208) translation initiation factor IF3 [Bacillus subtilis]	86 95	2.10E-54	562	729	g1770007	493-1	1218 Bt1Gc1562 Bt1G2365	
subtilis]			`.				*. *	
PROTEIN (GENERAL STRESS PROTEIN 22) [Bacillus	•		···		Ċ			
ALKYI		3.70E-66	673	671	g1703216	5104-5599	Bt1Gc1559	
(Z99111) ykrT [Bacillus subtilis]	37 100	8.90E-72	726	618	g2633727	4524-3295	1217 Bt1Gc1559 Bt1G2363	
INTERCENTO DECION [Bacillus subtilis]					(
coli] HYPOTHETICAL 38.9 KD PROTEIN IN DAT-SPO0E	49 100	6.40E-85	850	817	g3183453	3273-2215	1217 Bt1Gc1559 Bt1G2362	
FHUG [Bacillus subtilis] L-FUCULOSE PHOSPHATE ALDOLASE [Escherichia	37 100	1.80E-32	355	315	g120586	2173-1526	1217 Bt1Gc1559 Bt1G2361	
FERRIC	42 100	1.40E-50	526	736	g1706797	482-1483	1217 Bt1Gc1559 Bt1G2360	
				Score		*	NO	
NCBI gi description	% % Ident Cyrg	BlastP- d	BlastP Score	nap	NCBI gi	Position	D Contig Id Gene Id	
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(UROPORPHYRINOGEN III METHYLASE) (UROM) Racillus megaterium								
(UROGEN III METHYLASE) (SUMT)				٠				
UROPORPHYRIN-III C-METHYLTRANSFERASE	73	8 57	1.00E-38	414	479	g267052	8467-7948	1225 Bt1Gc1567 Bt1G2401
(AJ000974) YlnE protein [Bacillus subtilis]	100	7 30	1.10E-27	310	265	g2462961	7243-5768	1225 Bt1Gc1567 Bt1G2400
(AJ000974) YlnF protein [Bacillus subtilis]	99	8 33	6.00E-18	218	224	g2462962	6528-6055	1225 Bt1Gc1567 Bt1G2399
FACILITATOR) [Bacillus subtilis]								
NITRITE EXTRUSION PROTEIN (NITRITE	100	1 65	6.40E-101	1001	1391	g1171655	4011-5195	1225 Bt1Gc1567 Bt1G2398
[Bacillus subtilis]						-		
(Z99111) molybdopterin converting factor (subunit 1)	99	4 47	2.40E-14	184	184	g2633802	3700-3930	1225 Bt1Gc1567 Bt1G2397
[Bacillus subtilis]								
(Z99111) molybdopterin converting factor (subunit 2)	99	3 54	3.60E-43	456	423	g2633801	3287-3762	1225 Bt1Gc1567 Bt1G2396
subtilis]						•		•
(Z99111) molybdopterin biosynthesis protein [Bacillus	100	61	5.40E-136	1332	1328	g2633799	1947-3236	1225 Bt1Gc1567 Bt1G2395
subtilis]								
(Z99111) molybdopterin biosynthesis protein [Bacillus	100	6 51	1.30E-86	866	927	g2633798	890-1903	1225 Bt1Gc1567 Bt1G2394
NARA PROTEIN [Bacillus subtilis]	85	9 57	2.70E-79	797	848	g730103	1-869	1225 Bt1Gc1567 Bt1G2393
(AF080235) reductase homolog [Streptomyces cyanogenus]	8 2	5 47	2.80E-45	476	463		10332-10964	1224 Bt1Gc1568 Bt1G2392
family) [Bacillus subtilis]								•
(Z99107) similar to transcriptional regulator (Lrp/AsnC	99		8.20E-14	179	178	g2632968	10146-9892	1224 Bt1Gc1568 Bt1G2391
(AL021841) amiB [Mycobacterium tuberculosis]	100		1.50E-53	554	545	g2894215	8358-9549	1224 Bt1Gc1568 Bt1G2390
HYPOTHETICAL PROTEIN KIAA0083 [Homo sapiens]	18	2 27	5.50E-12	175	55	g2506893	6293-9003	1224 Bt1Gc1568 Bt1G2389
INTERGENIC REGION [Bacillus subtilis]								
HYPOTHETICAL 14.4 KD PROTEIN IN EPR-GALK	99	2 52	4.90E-32	351	339	g732333	5658-5278	1224 Bt1Gc1568 Bt1G2388
[Paenihacillus nolymyxa]								
[Bacillus thuringiensis] BACILLOLYSIN PRECURSOR (NEUTRAL PROTEASE)	53	9 33	1.10E-39	423	220	g128529	2264-5170	1223 Bt1Gc1569 Bt1G2387
transcriptional activator pick - Bacillus thuringiensis	<u>S</u>	8	7.80E-128	1255	1255	8212/289	3293-2441	1223 BUIGCISBY BUIG2386
thuringiensis]								1
hypothetical 5.3K protein - Bacillus thuringiensis [Bacillus	98	7 83	6.90E-17	208	208	g2127281	2348-2205	1223 Bt1Gc1569 Bt1G2385
(Z99110) similar to hypothetical proteins [Bacillus subtilis]	77	8 . 39	2.50E-108	1071	938	g2633696	1-2186	1223 Bt1Gc1569 Bt1G2384
[Bacillus subtilis]		. •		٠.				
respiratory nitrate reductase alpha chain - Bacillus subtilis	97	0 81	_	5206	5360	g2117582	7220-3656	1222 Bt1Gc1566 Bt1G2383
subtilis]								
ANAEROBIC REGULATORY PROTEIN [Bacillus	100) 56	1.70E-70	714	699 .	g1169717	81-791	1222 Bt1Gc1566 Bt1G2382
	S.A.	TOPIL	rron	Score	Score	,		
NCBI gi description	%	I %	BlastP-	BlastP	aat_	NCBI gi	Position	SEQ ID Contig Id Gene Id
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Bt1G	Bt1G	Bt1G	Bt1G	Bt1G	Bt1G	Bt1G	Bt1G		Bt1G	Bt1G	Bt1Gc15/2	BtiG	Bt1G	Bt1Gc1572	Bt1G	Bt1Gc1572		BtlG		Bt1G		Bt1G		₽. -1-0-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-	Bt1G	Bt1G	Bt1Gc1570	Contig Id	
Bt1Gc1574	Bt1Gc1574	Bt1Gc1574	Bt1Gc1574	Bt1Gc1574	Bt1Gc1574	Bt1Gc1574	Bt1Gc1574		Bt1Gc1574	Bt1Gc1572	01572	Bt1Gc1572	Bt1Gc1572	:1572	Bt1Gc1572	:1572		Bt1Gc1572	.•	Bt1Gc1572	•	Bt1Gc1570		R+10~1570	Bt1Gc1570	Bt1Gc1570	:1570	ig Id	: .
			Bt10								Bti	Btic	Bt10	Bt10	Bt10	Bt10				Bt10		Bt1(Bt10	Bt10	Bt10	Ge	
Bt1G2424	Bt1G2423	Bt1G2422	Bt1G2421	Bt1G2420	Bt1G2419	Bt1G2418	Bt1G2417		Bt1G2416	Bt1G2415	Bt1G2414	Bt1G2413	Bt1G2412	Bt1G2411	Bt1G2410	Bt1G2409		Bt1G2408		Bt1G2407		Bt1G2406	i	R+1G2405	Bt1G2404	Bt1G2403	Bt1G2402	Gene Id	
886	830	774	738	564	556	489	463	,	14	933	804	731	564	678	213	107		9.		2		375	i	213	209	997	303	Po	
8864-8301	8308-7751	7742-7389	7389-6631	5640-6458	5562-4966	4899-4331	4636-1999	į	142-405	9330-6993	8043-7318	7319-6021	5640-4477	6788-3625	2134-1841	1070-1448		958-17		281-1		3756-4556		3171-3744	2097-3014	997-2031	303-1016	Position	
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g3323048	g1730984	g1730985	g1730986	g1303795	g729154	g418361	g1303798		o1710749	g1169894	g2633716	g2633717	<u>g2635915</u>	g3721552	g4894353	<u></u> <u></u> <u> </u>		g3183273	j	g1881228		g1705428		01168677	g1168621	g4481748	g4481749	NCBI gi	
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196	609	436	541	815	320	2	185		250	77	565	321	265	209	142	297		412		159		713		2	1276	771	918	eo .	
239	611	368	598	763	365	661	1229	!	254	116	587	296	367	321	.173	340		508		173		561	ì	8	1231	761	922	BlastP Score	
3.6	1.4	7.7	3.3	1.1	1.6	6.9	4.40	ì	9	3.4	4.	. ພ ພ	9.8	5.4	3.5	7.1		1.1		2.4		2.7	;	y V	2.70	1.7	1.5	BlastP- Prob	1
3.60E-20	.40E-59	7.70E-34	3.30E-58	1.10E-75	1.60E-33	6.90E-65	4.40E-125	i	9 20F-22	3.40E-06	4.80E-57	3.30E-26	9.80E-34	5.40E-28	3.50E-13	7.10E-31	•	1.10E-48		2.40E-12		2.70E-54	:	5 60E-47	2.70E-125	1.70E-75	1.50E-92	8 %	
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(AE001246) conserved hypothetical protein	INTERGENIC REGION [Bacillus subtilis] HYPOTHETICAL 21.3 KD PROTEIN IN .	INTERGENIC REGION [Bacillus subtilis] HYPOTHETICAL 13.3 KD PROTEIN IN A	YPOI	(D8443)	COME	COME	(D8443)	subtilis]	GGAA [Bacillus subtilis]	MINOR	(Z99111 (Z91111)	(Z99111	(Z99121	(AB010958) xylanase [Clostridium thermocellum]	(AF065404) pXO1-138 [Bacillus anthracis]	(AF008)	[Bacillus subtilis]	PUTAT	AN AIF-DEFENDENI COVALENI BINDI TO THEIR SUBSTRATE. [Bacillus subtilis]	B001	UNDECAPRENOL KINASE) [Escherichia coli]	CIT	BCRB	BACITE ACINI TE ANGEORT BERMEACE DEOTEINI	CIT	(AF007865) BacS [Bacillus licheniformis]	(AF0078		
246)	HEI	HET	HET	2) ComED [Bacillus subtilis]	OPERON PROTEIN 1 [Bacillus subtilis]	OPERON PROTEIN 2 [Bacillus subtilis]	2) ComEC [Bacillus subtilis]	(l) sim	:) ykr		958):	404) ₁	220) YtcD [Bacillus subtilis]	s subt	IVE		488)	APR	RACI	Bacillus licheniformis	LPacillus lichenitormis]	ACI	865) I	865) BacR [Bacillus licheniformis]		
CONSE	CRE	ICAL	ICAL	mED	õ	õ	mEC		[Bacillus subtilis]	CHOI	ilar t) ykrI [Bacillus subtilis]	ilar t	xylan	0XO	YtcD	ilis]	NAD DAD		SIM	ENO	N RE	llus li	בין ש צוון	Z Z	BacS	BacR		
hed	GIOI , 21.3	, 13.3	, 28.3	Ва	PRO	PRO	[Bac		PRC	C AC	R N	cillu	o hyp	ase [-138	Bac		P)H	RA Z	LAR	LKI	SIST	cheni	A N.C.	ANS	[Bac	Вас	NC.	
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JENIC REGION [Bacillus subtilis] 246) conserved hypothetical protein [Treponema	INTERGENIC REGION [Bacillus subtilis] HYPOTHETICAL 21.3 KD PROTEIN IN AROD-COMER	INTERGENIC REGION [Bacillus subtilis] HYPOTHETICAL 13.3 KD PROTEIN IN AROD-COMER	HYPOTHETICAL 28.3 KD PROTEIN IN AROD-COMER					į		TEICHOIC ACIDS BIOSYNTHËSIS PROTEIN) similar to RNA polymerase sigma factor [Bacillus	ì) similar to hypothetical proteins [Bacillus subtilis]	二				ĒQ	AN AIP-DEPENDENT COVALENT BINDING OF AMP TO THEIR SUBSTRATE. [Bacillus subtilis]	(AB001488) SIMILAR TO ENZYMES WHICH ACT VIA		BACITRACIN RESISTANCE PROTEIN (PUTATIVE		Į Į	BACITRACIN TRANSPORT ATP-BINDING PROTEIN				
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Gene Id		Bt1G2448		Bt1G2449	B+1C2450	Bt1G2451				Bt1G2454		Bt1G2456		Bt1G2457				Bt1G2460	Bt1G2461	Bt1G2463
Position		6296-7237		6401-8806	0114-0086	1-721	1610 2597	4047-3367	7301-4674	1-1185	٠.	2787-1975		3199-4691	1 } } s	1001-1	; ; ;	4726-2369	6458-4749	7324-7070
NCBI gi		g3025110		g2493485	27404604	g1724006		87007761	g3122390	g1169253	· ·	g732340		g1881236)))	g2293322	•	g3914084	g1770035	g1770033
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BlastP Score	· -	520		1394	702	663	}	Ş	1110	1021		· 819	•	1645) }	8 33		2282	1733	236
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% Ident		38		54	<u>.</u>	69			37	57		60		2	1	54		63	65	58
Cyrg		100	:.:	100	3	52	3		78	90		100		68	I. ·	75	:	100	100	. 99
NCBI gi description	PROTEIN YDEX [Escherichia coli]	HYPOTHETICAL TRANSCRIPTIONAL REGULATOR IN HIPB-UXAB INTERGENIC REGION [Escherichia	coli]	HYPOTHETICAL SUGAR KINASE IN HIPB-UXAB	INTERGENIC REGION [Escherichia coli]	HYPOTHETICAL 49.0 KD PROTEIN IN CSPB-GLPP	INTERGENIC REGION [Bacillus subtilis]	UDPGlcNAc GlcNAc transferase [Streptococcus	<pre>pneumoniae] 5-METHYLTETRAHYDROFOLATEHOMOCYSTEI</pre>	ANAEROBIC C4-DICARBOXYLATE TRANSPORTE	PYROPHOSPHORYLASE) (TMP-PPASE) (THIAMIN-PHOSPHATE SYNTHASE) [Bacillus subtilis]	HYDROXYETHYLTHIAZOLE KINASE (4-METHYL	BETA-HYDROXYETHYLTHIAZOLE KINASE) (THZ KINASE) /TH KINASE) IBacillus subtilis]	(AB001488) PROBABLE DNA TOPOISOMERASE III	[Bacillus subtilis]	(AF008220) branch-chain amino acid transporter [Bacil subtilis]	INTERGENIC REGION [Escherichia coli]	MUTS2 PROTEIN [Bacillus subtilis]	(Z75208) hypothetical protein [Bacillus subtilis]	(275208) hypothetical protein [Bacillus subtilis]
Score Bt1Gc1575 Bt1G2448 6296-7237 g3025110 455 520 6.00E-50 38 100 Bt1Gc1575 Bt1G2449 6401-8806 g2493485 1432 1394 1.50E-142 54 100 Bt1Gc1575 Bt1G2450 0114-0086 g2493485 507 586 6 10E-57 41 100	Bt1Gc1575 Bt1G2448 6296-7237 g3025110 455 520 6.00E-50 38 100 Bt1Gc1575 Bt1G2449 6401-8806 g2493485 1432 1394 1.50E-142 54 100 Bt1Gc1575 Bt1G2450 0114-0086 g2495606 507 586 6 10E-57 41 100	Bt1Gc1575 Bt1G2449 6401-8806 g2493485 1432 1394 1.50E-142 54 100	Bt1Gc1575 Bt1G2449 6401-8806 g2493485 1432 1394 1.50E-142 54 100 Bt1Gc1575 Bt1G2450 0114-0086 g2493485 507 586 6 10E-57 41 100	R+1Gc1575 R+1G2450 0114-0086 02405606 507 586 6 10F-57 41 100			Bt1G2451 1-721 g1724006 827 663 4.20E-65 69 52	Bt1Gc1576 Bt1G2451 1-721 g1724006 827 663 4.20E-65 69 52	Bt1Gc1576 Bt1G2451 1-721 g1724006 827 663 4.20E-65 69 52 Bt1Gc1576 Bt1G2452 4649-3587 g4009481 608 604 7.50E-59 40 100	Bt1Gc1576 Bt1G2451 1-721 g1724006 827 663 4.20E-65 69 52 Bt1Gc1576 Bt1G2452 4649-3587 g4009481 608 604 7.50E-59 40 100 Bt1Gc1576 Bt1G2453 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4726-2369 g3914084 2531 2282 1.20E-236 63 100 Bt1Gc1580 Bt1G2462 7059-6526 g1770035 1974 1733 1.70E-178 65 100 Bt1Gc1580 Bt1G2462 7059-6526 g1770034 387 352 3.80E-32 45 99	Bt1Gc1577 Bt1G2455 1962-1297 g732341 665 592 1.40E-57 59 100 Bt1Gc1577 Bt1G2456 2787-1975 g732340 804 819 1.20E-81 60 100 Bt1Gc1577 Bt1G2457 3199-4691 g1881236 1658 1645 3.70E-169 64 68 Bt1Gc1580 Bt1G2458 1001-1 g2293322 888 833 4.10E-83 54 75 Bt1Gc1580 Bt1G2459 2188-323 g418534 554 603 9.60E-59 48 89 Bt1Gc1580 Bt1G2460 4726-2369 g3914084 2531 2282 1.20E-236 63 100 Bt1Gc1580 Bt1G2461 6458-4749 g1770035 1974 1733 1.70E-178 65 100 Bt1Gc1580 Bt1G2462 7059-6526 g1770034 387 352 3.80E-32 45 99	Bt1Gc1577 Bt1G2456 2787-1975 g732340 804 819 1.20E-81 60 100 Bt1Gc1577 Bt1G2457 3199-4691 g1881236 1658 1645 3.70E-169 64 68 Bt1Gc1580 Bt1G2458 1001-1 g2293322 888 833 4.10E-83 54 75 Bt1Gc1580 Bt1G2459 2188-323 g418534 554 603 9.60E-59 48 89 Bt1Gc1580 Bt1G2460 4726-2369 g3914084 2531 2282 1.20E-236 63 100 Bt1Gc1580 Bt1G2461 6458-4749 g1770035 1974 1733 1.70E-178 65 100 Bt1Gc1580 Bt1G2462 7059-6526 g1770034 387 352 3.80E-32 45 99	Bt1Gc1577 Bt1G2457 3199-4691 g1881236 1658 1645 3.70E-169 64 68 Bt1Gc1580 Bt1G2458 1001-1 g2293322 888 833 4.10E-83 54 75 Bt1Gc1580 Bt1G2459 2188-323 g418534 554 603 9.60E-59 48 89 Bt1Gc1580 Bt1G2460 4726-2369 g3914084 2531 2282 1.20E-236 63 100 Bt1Gc1580 Bt1G2461 6458-4749 g1770035 1974 1733 1.70E-178 65 100 Bt1Gc1580 Bt1G2462 7059-6526 g1770034 387 352 3.80E-32 45 99	Bt1Gc1577 Bt1G2457 3199-4691 g1881236 1658 1645 3.70E-169 64 68 Bt1Gc1580 Bt1G2458 1001-1 g2293322 888 833 4.10E-83 54 75 Bt1Gc1580 Bt1G2459 2188-323 g418534 554 603 9.60E-59 48 89 Bt1Gc1580 Bt1G2460 4726-2369 g3914084 2531 2282 1.20E-236 63 100 Bt1Gc1580 Bt1G2461 6458-4749 g1770035 1974 1733 1.70E-178 65 100 Bt1Gc1580 Bt1G2462 7059-6526 g1770034 387 352 3.80E-32 45 99	Bt1Gc1580 Bt1G2458 1001-1 g2293322 888 833 4.10E-83 54 75 Bt1Gc1580 Bt1G2459 2188-323 g418534 554 603 9.60E-59 48 89 Bt1Gc1580 Bt1G2460 4726-2369 g3914084 2531 2282 1.20E-236 63 100 Bt1Gc1580 Bt1G2461 6458-4749 g1770035 1974 1733 1.70E-178 65 100 Bt1Gc1580 Bt1G2462 7059-6526 g1770034 387 352 3.80E-32 45 99	Bt1Gc1580 Bt1G2458 1001-1 g2293322 888 833 4.10E-83 54 75 Bt1Gc1580 Bt1G2459 2188-323 g418534 554 603 9.60E-59 48 89 Bt1Gc1580 Bt1G2460 4726-2369 g3914084 2531 2282 1.20E-236 63 100 Bt1Gc1580 Bt1G2461 6458-4749 g1770035 1974 1733 1.70E-178 65 100 Bt1Gc1580 Bt1G2462 7059-6526 g1770034 387 352 3.80E-32 45 99	Bt1Gc1580 Bt1G2459 2188-323 g418534 554 603 9.60E-59 48 89 Bt1Gc1580 Bt1G2460 4726-2369 g3914084 2531 2282 1.20E-236 63 100 Bt1Gc1580 Bt1G2461 6458-4749 g1770035 1974 1733 1.70E-178 65 100 Bt1Gc1580 Bt1G2462 7059-6526 g1770034 387 352 3.80E-32 45 99	Bt1Gc1580 Bt1G2460 4726-2369 g3914084 2531 2282 1.20E-236 63 100 Bt1Gc1580 Bt1G2461 6458-4749 g1770035 1974 1733 1.70E-178 65 100 Bt1Gc1580 Bt1G2462 7059-6526 g1770034 387 352 3.80E-32 45 99	Bt1Gc1580 Bt1G2461 6458-4749 g1770035 1974 1733 1.70E-178 65 100 Bt1Gc1580 Bt1G2462 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	75 1200 1244	0 418 000	906	0 1344 1350	72 3115 2861		4 815 829		٠.	392	51 376 390	49 1419 1442	919		162	43 649 671	-	2459	1341 1	05 1005 946		 22 853 806	100	5 250 201	5 344 347	•		7.0	aat_		
	1.10E-126	2.30E-04		6.70E-138	5.10E-298		1.10E-82				3.60E-36	1.20E-147			4.40E-15	6.00E-66				4.30E-95	·	3.00E-80		3 10E_35	1.30E-31		9.70E-27	rrop Ident		Table 1	:
·	48 100	38 72	_	51 100	69 100	٠.	58 100			_	59 99	74 100	68 100	٠,٠	32 .99	67 40				62 100		70 100	1	ა	79 99		67 99	ar cyrg	%		•.
(GOLD) (AEGETATIAE LUGIETA II) (AEGIT) [Dacillas		HE) ALKALINE PHOSPHATASE SYNTHESIS SENSOR) (AF008220) DNA-polymerase I [Bacillus subtilis]	(FAPY-DNA GLYCOSYLASE) [Bacillus firmus]		subtilis]			aerogenes] (AB001488) FUNCTION UNKNOWN. [Bacillus subtilis]	(AB0) (AF008220) 3-ketoacyl-ACP reductase [Bacillus subtilis]	thermoautotrophicum]) (AF008220) arginine succinate lyase [Bacillus subtilis]	[Entamoeba histolytica]				(MTA/SAH NUCLEOSIDASE) (P46) [Bacillus subtilis]			INTERGENIC REGION [Bacillus subtilis]			HYPOTHETICAL 11.4 KD PROTEIN IN MFD-DIVIC		NCBI gi description		· · · · · · · · · · · · · · · · · · ·

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	46 BtlGc1587	1246 Bt1Gc1587 1	1246 Bt1Gc1587 1	1245 Bt1Gc1589 1	1245 Bt1Gc1589 I		1245 Bt1Gc1589 I		1245 BilliGe1589 I		1245 Bt1Gc1589 I	1244 Bt1Gc1588 1	1244 Bt1Gc1588 1	Bt1Gc1588	1243 Bt1Gc1591 I	1	1243 Bt1Gc1591 I	1243 Bt1Gc1591 I	. ·	2	Bt1Gc1591	2 Bt1Gc1584	ID Contig Id	
	Bt1G2519	Bt1G2518	Bt1G2517	Bt1G2516	Bt1G2515	٠	Bt1G2514		Bt1G2513		Bt1G2512	Bt1G2511	Bt1G2510	Bt1G2509	Bt1G2508		Bt1G2507	Bt1G2506			Bt1G2505	Bt1G2504	Gene Id	
	5444-4022	1176-3239	1-366	4655-5626	4533-3070		2965-1652		1485-604		582-1	5797-4670	1675-4626	Jan-79	4439-5326		3618-3199	3106-2783			1475-404	5257-7245	Position	
i i	g461637	g124464	g141471	g586856	g2127114		g585034		g586857		g586858	g1881236	g538903	g4098489	g2507254	- !	g732301	g732304			g728897	g2507484	NCBI gi	
	168	2632	464	652	2029		1165		1291		659	1183	997	541	83	,	355	268	•	***	1410		nap Score	
	276	2519	417	651	2033		1162		1232		626	1161	1208	567	208	. !	244	255			1189	1771	BlastP Score	
	4.30E-24	8.90E-262	4.90E-39	7.90E-64	2.80E-210		5.60E-118		2.10E-125		3.50E-61	7.10E-118	7.40E-123	6.30E-55	2.70E-16		1.10E-20	7.20E-22			7.70E-121	1.60E-182	BlastP- Prob	
	23	70	72	45	79		54		86	.	68	63	32	53	39	,	50	4			77	57	% Ident (
	100	100	. 65	100	100		100	. •	0		% -	52	100	56	30		99	99				100	% Cvrg	
	EFFLUX TRANSPORTER 1) [Bacillus subtilis]	PHOSPHOPROTEIN ORFU) (CSI9) [Bacillus subtilis] IMMUNE INHIBITOR A PRECURSOR [Bacillus	INTERGENIC REGION [Bacillus subtilis] TRANSALDOLASE-LIKE PROTEIN (20 KD	HYPOTHETICAL 38.0 KD PROTEIN IN GIRA-GUAB	IMP dehydrogenase - Bacillus subtilis [Bacillus subtilis]	CARBOXYPEPTIDASE) (CPASE) (PBP5) [Bacillus	D-ALANYL-D-ALANINE CARBOXYPEPTIDASE	PROTEIN 7) (SOI7) [Bacillus subtilis]	JUTERGENIC REGION (SUPEROXIDE-INDUCIBLE	INTERGENIC REGION [Bacillus subtilis]	[bacinus suonns] HYPOTHETICAL 21.4 KD PROTEIN IN DACA-SERS	(ABO) (1488) PROBABLE DNA TOPOISOMERASE III	phage infection protein precursor - Lactococcus lactis subsp.	(U78600) putative ptsG protein [Streptococcus mutans]	AMINOPEPTIDASE AMPS [Bacillus subtilis]	INTERGENIC REGION (ORF1) [Bacillus subtilis]	INTERGENIC REGION (ORF2) (ORF3) [Bacillus subtilis] HYPOTHETICAL 15.7 KD PROTEIN IN MURC-AROA	HYPOTHETICAL 12.4 KD PROTEIN IN MURC-AROA	SYNTHETASE) (3-DEOXY-D-ARABINO- HEPTULOSONATE 7-PHOSPHATE SYNTHASE) / CHORISMATE MUTASE [Bacillus subtilis]	ALDOLASE (PHOSPHO-2-KETO-3- DEOXYHEPTONATE ALDOLASE) (DAHP	PHOSPHO-2-DEHYDRO-3-DEOXYHEPTONATE	TRANSKETOLASE [Bacillus subtilis]	NCBI gi description	. •

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1249 Bt1Gc1592 Bt1G2534	1248 Bt1Gc1590 Bt1G2533 10	1248 Bt1Gc1590 Bt1G2531 7: 1248 Bt1Gc1590 Bt1G2532 8:		1248 Bt1Gc1590 Bt1G2530 49		1248 Bt1Gc1590 Bt1G2529 34		D11 01040	1248 B+1G-1500 B+1G2528 24			Bt1G2527	1247 Bt1Gc1594 Bt1G2526 51	1247 Bt1Gc1594 Bt1G2525 41	1247 Bt1Gc1594 Bt1G2524 33	1247 Bt1Gc1594 Bt1G2523 22	1247 BUIGC1394 BUIGZ3ZZ I	Bt1Gc1587 Bt1G2521		1246 Bt1Gc1587 Bt1G2520 54	SEQ ID Contig Id Gene Id I NO
1-1080	10381-8367	7234-6137 8168-7263		4938-3520		3480-2491			2474-1404			1475-159	5161-6451	4169-5224	3387-4166	2274-1840	C24-02/1	5650-6699		5456-4265	Position
g584749	g1731060	g1256412 g1709891		g1706441	•	g585606		800000	a585607	• .		g585605	g1651978	g421277	g1197640	g2226212	6/010/0	g2522016		g728970	NCBI gi
1048	2059	1868 819	•	1795 . 1743	٠.,٠	1423		5	1540		14 -	1420	910	. 843	876	204	1040	1099	· •	180	nap s
883	2038	1720 788		1743	*	1286		1	1540	,,,,,,,		1365	903	901	893	194	1170	1107		293	BlastP Score
8.90E-98	8.30E-211	4.10E-177 2.40E-78		1.50E-179		4.00E-131		4.70E-100	4 00E-158			1.70E-139	1.60E-90	2.50E-90	1.80E-89	2.10E-15	0.005-121	3.80E-112		6.80E-26	BlastP- Prob
57	61	100 59		73		80		5	80			71	45	4	61	36	7	2 2		21	%
98	97	. 10 E		100		100			3				<u> </u>	100	100	99	5			100	% Cvrg
BFMBAA INTERGENIC REGION [Bacillus subtilis] N-ACYL-L-AMINO ACID AMIDOHYDROLASE	(PHOSPHOI KANSBOITKY LASE) [Bacillus Subtilis] PUTATIVE SIGMA L-DEPENDENT TRANSCRIPTIONAL BEGLII ATOR IN MMCE	(US1099) leucine dehydrogenase LeuDH [Bacillus cereus] PROBABLE PHOSPHATE BUTYRYLTRANSFERASE	BRANCHED-CHAIN ALPHA-KETO ACID DEHYDROGENASE COMPLEX (E3) (DIHYDROLIPOAMIDE DEHYDROGENASE) (LPD-VAL) []	(E1)) (BCKDH E1-ALPHA) [Bacillus subtilis] LIPOAMIDE DEHYDROGENASE COMPONENT OF	SUBUNIT (BRANCHED-CHAIN ALPHA-KETO ACID DEHYDROGEÑASE COMPONENT ALPHA CHAIN	2-OXOISOVALERATE DEHYDROGENASE ALPHA	DEHYDROGENASE COMPONENT BETA CHAIN (E1))	SUBUNIT (BRANCHED-CHAIN ALPHA-KETO ACID	TRANSACYLASE) [Bacillus subtilis]	(DIHYDROLIPOAMIDE BRANCHED CHAIN	BRANCHED-CHAIN ALPHA-KETO ACID	LIPOAMIDE ACYLTRANSFERASE COMPONENT OF	[Yersınıa pseudotuberculosis] (D90901) hypothetical protein [Synechocystis sp.]	CDP-glucose-4,6-dehydratase - Yersinia pseudotuberculosis	(U46859) DdhA [Yersinia enterocolitica (type 0:8)]	(Y14082) hypothetical protein [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis]	(AB007638) dehydrogenase [Bacillus subtilis]	EFFLUX TRANSPORTER 2) [Bacillus subtilis]	MULTIDRUG RESISTANCE PROTEIN 2 (MULTIDRUG-	NCBI gi description

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1254 Bt10	1254 Bt10	1254 Bt10	1254 Bt10	1254 Bt10	1254 Bt10	1253 Bt10	1253 Bt10	1253 Bt10	1253 Bt10		1253 Bt10	1253 Bt10	1253 Bt10	1253 Bt10		.1252 Bt10	1252 Bt10			1251 Rt1Gc1505	1251 Bt1Gc1595		SEQ Cor
Bt1Gc1600	Bt1Gc1600	Bt1Gc1600	Bt1Gc1600	Bt1Gc1600	Bt1Gc1600	Bt1Gc1596	Bt1Gc1596	Bt1Gc1596	Bt1Gc1596		Bt1Gc1596	Bt1Gc1596	Bt1Gc1596	Bt1Gc1596		Bt1Gc1598	Bt1Gc1598	-					Contig Id
Bt1G2568	Bt1G2567	Bt1G2566	Bt1G2565	Bt1G2564	Bt1G2563	Bt1G2562	Bt1G2561	Bt1G2560	Bt1G2559		Bt1G2558	Bt1G2557	Bt1G2556	Bt1G2555		Bt1G2554	Bt1G2553			R+1G2552	Bt1G2551		Gene Id
3827-2964	2918-1250	1339-1034	1019-684	671-390	1362-1	8381-9274	7288-7900	6784-6638	4846-6075		4232-3039	2566-3973	759-1841	1-757		1472-2907	2640-683		70.0	7610-4504	4501-3725		Position
g134771	g606187	g132771	g141394	'g132830	g134740	g2506131	g2507080	g1621448	g1731082		g3258413	g1177029	g2226255	g2226256	(g136144	g1176954		6110020	0115626	g1176556	•	NCBI gi
744	576	422	283	433	147	1207	275	123	942		305	96	1014	630.		2449	688		1000	4086	837		nap Score
729	528	342	286	433	216	1152	342	134	931		403	188	984	633		2407	735	•	1001	4004	782	٠	BlastP Score
4.30E-72	6.30E-64	4.40E-31	3.80E-25	9.90E-41	9.80E-18	6.40E-117	4.40E-31	4.80E-09	1.70E-93		1.50E-37	4.30E-14	4.10E-99	6.40E-62		6.60E-250	9.90E-73			-	1.00E-77	-	BlastP- Prob
48	34	80	49	. 60	40	79	33	56	46		29	41	56	51		99	56			77	60		% % Ident Cvrg
100	100	99	99	99	66	33	,100	98	100		100	20	100	51		100	74			07	100		%Cvrg
[Escherichia coli] STAGE IV SPORULATION PROTEIN FB [Bacillus	subtilis] (U18997) ORF_f495; orfF of ECMRED, uses 2nd start	INTERGENIC REGION (ORF X) [Bacillus subtilis] 50S RIBOSOMAL PROTEIN L21 (BL20) [Bacillus	subtilis] HYPOTHETICAL 12.3 KD PROTEIN IN RPLU-RPMA	50S RIBOSOMAL PROTEIN L27 (BL30) (BL24) [Bacillus	(ACONITASE) [Bacillus subtilis] STAGE 0 SPORULATION PROTEIN B [Bacillus subtilis]	(NODULATION PROTEIN B) [Rhizobium sp. NGR234] ACONITATE HYDRATASE (CITRATE HYDRO-LYASE)	CHITOOLIGOSACCHARIDE DEACETYLASE	(U72073) Cotl. [Bacillus subtilis]	HYPOTHETICAL 44.7 KD PROTEIN IN GLNQ-ANSR	synthetase [Pyrococcus horikoshii]	SREGION [] (AP000007) 404aa long hypothetical alanyl-tRNA	HYPOTHETICAL 45.4 KD PROTEIN IN THIAMINASE I	(Y14084) hypothetical protein [Bacillus subtilis]	(Y14084) hypothetical protein [Bacillus subtilis]	EI EMENT IS231C [Racillus thuringiensis]	5'REGION [Bacillus subtilis] TRANSPOSASE FOR INSERTION SEOUENCE	HYPOTHETICAL 36.9 KD PROTEIN IN ACDA	(CARBAMOYL-PHOSPHATE SYNTHETASE	DADIMILIME CAECIEIC I VACE CHVIM	INTERGENIC REGION (ORF2) [Bacillus caldolyticus] CARRAMOVI _DHOCDHATE SVNTHASE	HYPOTHETICAL 27.6 KD PROTEIN IN PYRAB-PYRD	(DIHYDROOROTATE OXIDASE) (DHODEHASE)	NCBI gi description

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	NO County In	1254 Bt1Gc1600	1254 Bt1Gc1600	1254 Bt1Gc1600	1254 Bt1Gc1600		1254 Bt1Gc1600	1254 Bt1Gc1600			1255 Bt1Gc1597			1255 Bt1Gc1597	1255 Bt1Gc1597	1255 Bt1Gc1597			1255 Bt1Gc1597	1256 Bt1Gc1602	1256 Bt1Gc1602	1256 Bt1Gc1602	1256 Bt1Gc1602
	Cene IO) Bt1G2569) Bt1G2570	Bt1G2571	Br1G2572) Bt1G2573) Bt1G2574			/ Bt1G25/6	•		7 Bt1G2580	7 Bt1G2581	7 Bt1G2582			7 Bt1G2584	2 Bt1G2585	2 Bt1G2586	2 Bt1G2587	2 Bt1G2588
5 } } !	rosition	4425-3823	5497-4700	6186-5506	6755-6240	0700-0240	7621-6773	7989-7678	-	349-927	1736 1374	2325-1960	3078-873	3406-4686	6392-2362	7117-6539		10276-7909	10276-9732	811-257	1891-836	2339-1894	3399-2398
	NCDI 61	g134770	g400260	g400259	9266570	6,000	g266569	g266568		g1405459	g1405458	91149666	g1787702	g586861	g80261	e2634185	C	g1652519	g2506131	g1706593	g1731048	g1703004	g1731047
aat_	nap Score	387	1015	650	307	0	623	407		23.2 24.2 25.2 26.2 26.2 26.2 26.2 26.2 26.2 26	207	433	322	1120	221	346		317	756	721	1223	443	269
BlastP	Score	423	875	668	8	102	606	307	! !) 1 1 1 1	406	433	275	1069	292	422	-	350	771	721	1130	392	245
BlastP-	Prob	1.10E-39	1.40E-87	1.20E-65	3 90E-14	J.70E-14	4.60E-59	2.20E-27		1.50E-53	3.40E-24	9 90E-41	8.40E-37	4.00E-108	2.70E-24	1.50E-39		2.50E-31	1.50E-76	3.00E-71	1.40E-114	2.20E-36	8.30E-21
%	Ident Cyrg	48	71	60	3	ູ	46	79	- .	6 6	7 0	3 4	31	49	37	45	i	25	80	74	65	59	34
%	Cyrg	77	100	100	6	77	100	ω	;	3 4	9 %	و ر	98	. 100	13	99	;	83	20	99	100	99	.99
	g INCDI Bi description,	subtilis] 7 STAGE IV SPORULATION PROTEIN FA [Bacillus	subtilis] 0 SEPTUM SITE-DETERMINING PROTEIN MIND	[Bacillus subtilis] 0 SEPTUM SITE-DETERMINING PROTEIN MINC		subtilis		stearothermophilus] 1 ROD SHAPE-DETERMINING PROTEIN MREB [Bacillus	subtilis	(27323		9 (X86498) IS1136 DNA [Clostridium perfringens]		0 HYPOTHETICAL 48.6 KD PROTEIN IN SERS-DNAZ INTERGENIC REGION [Bacillus subtilis]		9 (Z99113) similar to thiol:disulfide interchange protein		(D9090	0 ACONITATE HYDRATASE (CITRATE HYDRO-LYASE) (ACONITASE) [Bacillus subtilis]	9 ELONGATION FACTOR P (EF-P) [Bacillus subtilis]	0 PUTATIVE PEPTIDASE IN GCVT-SPOIIIAA INTERGENIC REGION [Bacillus subtilis]		9 HYPOTHETICAL 20.2 KD PROTEIN IN GCVT- SPOIIIAA INTERGENIC REGION [Bacillus subtilis]

Table 1

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Bt1Gc1607	1259 Bt1Gc1607 B	1258 Bt1Gc1599 B	1258 Bt1Gc1599 B	1258 Bt1Gc1599 B		1258 Bt1Gc1599 B	1258 Bt1Gc1599 B	1258 Bt1Gc1599 B	•	1258 Bt1Gc1599 B	1257 Bt1Gc1603 B	1257 Bt1Gc1603 B	1257 Bt1Gc1603 B		1257 Bt1Gc1603 B	1257 Bt1Gc1603 B	1257 Bt1Gc1603 B		1257 Bt1Gc1603 B	٠	1257 Bt1Gc1603 B	1257 Bt1Gc1603 B	• .	1256 Bt1Gc1602 B	1256 Bt1Gc1602 B	•	1256 Bt1Gc1602 B		1256 Bt1Gc1602 B		B+1C+1602	700 BITCC1007 B	C 0410 -1600	ID Contig Id		•
Bt1G2613	Bt1G2612	Bt1G2611	Bt1G2610	Bt1G2609		Bt1G2608	Bt1G2607	Bt1G2606		Bt1G2604	Bt1G2603	Bt1G2602	Bt1G2601		Bt1G2600	Bt1G2599	Bt1G2598		Bt1G2597		Bt1G2596	Bt1G2595		Bt1G2594	Bt1G2593		Bt1G2592		Br1G2591	. 06670119	0+163400	Bt102589	2177500	Gene Id		
2579-982	1022-1	9821-8858	8861-7224	6737-7202		7761-5353	6023-3521	3302-2649		791-1	9134-8087	8066-7551	5722-6966		5264-3281	5620-3894	3271-3816		2514-2187		942-2141	137-721		6232-6574	7536-5816		5668-5243		5188-4532	C7C+-C71+	1106 1603	3122-4073	100 1070	Position		
g1805440	g80410	g2634163	g2634164	g1945649		g4678633	g1652657	g1817539	٠	g1881327	g1060858	g2626825	g2126617		g2648784	g2226188	g730008		g1622733		g728970	g1945679		g3582220	g586030	· :	g1731043	ď	01724017	81/31043	~1721046	g1/31046	1701040	NCBI gi		
872	214	263	1184	119		202	645	163		862	773	609	522		131	524	325		200		1488	350		246	104		59		448	155	.	1087	2	nap Score	994	
	•																	÷	٠.	•			•			-	599						-	BlastP Score		
620	269	326	1055	186		279	661	227		742	820	615	508		248	684	363		233		1194	375		266	137		599		457	155	5	080		st P		_
1.50E-60	2.40E-23	2.20E-29	1.20E-106	1.50E-14		7.50E-24	6.90E-65	6.70E-19		1.80E-73	9.70E-82	5.10E-60	1.10E-48		2.20E-20	2.50E-67	2.60E-33		1.60E-19		2.30E-121	1.40E-34		4.90E-23	1.10E-08		2.50E-58		2 80F_43	3.0UE-09	3 80E 00	8.10E-110		BlastP- Prob		Table 1
37	25	39	43	30		24	46	28		67	46	68	31		29	29	38		43		72	39		43	40		79	:	44	26	3	8	`	% % Ident Cvrg	· .	
97	88	75	100	94		79	57	99		S 2.	52	99	100		39	00 <u>1</u>	99		99		100	99		99	3. 38		99	į	<u>.</u>	. 77	3	100		%Cvrg		
(D50453		(Z99113)	_	_	[Streptomyces coelicolor]	_) (D78508) YfiO [Bacillus subtilis]	subtilis]	(AB001488) SIMILAR TO YDDS_BACSU. [Bacillus		D83967) YfkM [Bacillus subtilis]		protein (dppA) [Archaeoglobus fulgidus]			MODULATOR OF DRUG ACTIVITY B [Escherichia coli]	fibrisolvens]	_	EFFLUX TRANSPORTER 2) [Bacillus subtilis]		(Z94043) hypothetical protein [Bacillus subtilis]	lactis]	(AE001)		SPOIIIAA INTERGENIC REGION [Bacillus subtilis]			-	SPOIIIAA INTERGENIC REGION [Bacillus subtilis]				NCBI gi description		

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	1263 Bt1Gc1608	1263 Bt1Gc1608	1262 Bt1Gc1606	1262 Bt1Gc1606	1262 Bt1Gc1606	1262 Bt1Gc1606	1262 Bt1Gc1606	1261 Bt1Gc1609	1261 Bt1Gc1609	1260 Bt1Gc1604	1260 Bt1Gc1604	1260 Bt1Gc1604		1260 Bt1Gc1604	1260 Bt1Gc1604	1260 Bt1Gc1604	1259 Bt1Gc1607	1259 Bt1Gc1607	1259 Bt1Gc1607	1259 Bt1Gc1607	1259 Bt1Gc1607	SEQ ID Contig Id NO
	8091	809	1606	1606	1606	1606	906	1609	609	604	604	604		604	604	604	607	607	607	607	607	J Id
	Bt1G2633	Bt1G2632	Bt1G2631	Bt1G2630	Bt1G2629	Bt1G2628	Bt1G2627	Bt1G2626	Bt1G2625	Bt1G2624	Bt1G2623	Bt1G2622		Bt1G2621	Bt1G2620	Bt1G2619	Bt1G2618	Bt1G2617	Bt1G2616	Bt1G2615	Bt1G2614	Gene Id
	3826-2191	2196-1	8888-9454	6922-8903	5775-6779	4723-5745	3025-4329	4989-5863	766-4931	6167-5769	5722-5036	5022-4570		4579-3547	1146-3077	964-317	7569-8191	6761-7552	6403-6768	5805-6284	2849-5786	Position
	g585920	g1495275	g4589058	g2120367	g421459	g465642	g133292	g585251	g2623773	g3323187	g1945108	g1945109	000	93025119	g3025120	g1945113	g730103	g730100	g3915538	g3915537	g2635168	NCBI gi
	2484	3744	751	1748	1386	1113	835	260	966	282	738	442		1337	2151	836	646	731	428	491	4439	aat_ nap Score
	2397	3612	778	1801	1126	950	855	. 380	1152	292	738	449	į	1299	1937	806	659	755	434	441	4352	BlastP Score
	7.50E-249	0	2.70E-77	1.10E-187	3.60E-114	1.60E-95	1.90E-85	4.10E-35	9.50E-115	8.70E-26	4.80E-73	2.00E-42		1 70E-132	2.10E-208	3.00E-80	1.10E-64	7.50E-75	7.80E-41	1.40E-41	. 0	BlastP- Prob
	∞	100	77	55	<u>∞</u>	65	42	31	37	43	61	52	į	75	66	76	60	55	66	57	&	% Ident
-	46	69	37	100	100	100	100	· 58	11	99	100	99		3	100	100	61	100	. 99	99 .	100	% Cyrg
	anthracis] 5 DNA-DIRECTED RNA POLYMERASE BETA CHAIN	phosphoglycerate mutase [Bacillus stearothermophilus] (X89230) DNA-directed RNA polymerase [Bacillus			IN GAP 5'REGION (ORF1) [Bacillus megaterium] glyceraldehyde-3-phosphate dehydrogenase (EC 1.2.1.12) -			3 POLYKETIDE SYNTHASE HETM [Anabaena sp.]	pallidum] [(AF004835) tyrocidine synthetase 3 [Brevibacillus brevis]	[Bacillus subtilis] (AE001257) conserved hypothetical protein [Treponema			INTER	PROTEIN YDIF [Bacillus subtilis]		(D8880	NARA	IN I EKGENIC KEGION [Bacillus subtilis] NARQ PROTEIN [Bacillus subtilis]				NCBI gi description

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INTERGENIC REGION [Bacillus subtilis]	86 100 100		2137 ^(r) 2 70E-221)	a1730257	221-768	1 B+1G2652	1266 B+1Gc1611	3
initiation point for translation of NPPase. [Homo sapiens] HYPOTHETICAL 47.3 KD PROTEIN IN WAPA-LICT	58 36	3.70E-34 5		. 371	501	g1177011	7165-7623	0 Bt1G2651	1265 Bt1Gc1610	12
nucleo	* 43.			-						
(D12485) The first in-frame ATG codon is located at	27 23	1.20E-19 2		1 246	101	g219944	4116-7623	0 Bt1G2650	1265 Bt1Gc1610	12
(Z99107) yesK [Bacillus subtilis]		3.50E-06 2			116	g2633006	5691-5307	0 Bt1G2649	1265 Bt1Gc1610	12
burgdorferi]				<i>;</i> .						
(AE001145) prolyl-tRNA synthetase (proS) [Borrelia	48 100		7 2.90E-114	1127	1163	g2688299	4938-3523	0 Bt1G2648	1265 Bt1Gc1610	12
(OBEA) [Classidium postinional	36 99	7.80E-25 3		2 283	292	g141085	1924-1385	0 Bt1G2646	65 Bt1Gc1610	1265
•					649	g2415740	1256-663			1265
_		-			708	g2415741	632-54			1265
(Z99104) ybaF [Bacillus subtilis]	68 72	1.70E-52 6		544	669	g2632414	7814-7244	5 Bt1G2643	64 Bt1Gc1605	1264
HYDROLYASE) [Bacillus subtilis]				٠.					٠	
(PSEUDOURIDINE SYNTHASE I) (URACIL										•
IRNA PSEUDOURIDINE SYNTHASE A	00 TOO	2.40E-69 S 5		\$ 703	817	g3915172	/223-6486	5 Bt1G2642	1264 Bt1Gc1605	12
50S RIBOSOMAL PROTEIN L13 [Bacillus subtilis]					621	g2500252	6330-5896			1264
30S RIBOSOMAL PROTEIN S9 (BS10) [Bacillus subtilis]					548	g2507325	5871-5482			12
5'REGION (ORF1) [Bacillus subtilis]		•								
(AUTOLYSIN) [Bacillus subtilis] HYPOTHETICAL 17.6 KD PROTEIN IN CWLD	29 99	7.10E-15 2		1 189	204	g1723295	5316-4891	5 Bt1G2639	1264 Bt1Gc1605	12
ALANINE AMIDASE (CELL WALL HYDROLASE)		•								
GERMINATION-SPECIFIC N-ACETYLMURAMOYL-L-	59 100	1.90E-60 5		7 619	727	g1706210	4824-4111	5 Bt1G2638	1264 Bt1Gc1605	12
INTERGENIC REGION [Bacillus subtilis]						c				
[Bacillus subtilis] HYPOTHETICAL 38.6 KD PROTEIN IN CWLD-GERD	66 100	1.20E-113 6		3 1121	1238	g1723296	3963-2902	5 Bt1G2637	64 Bt1Gc1605	,126 4
[Bacillus subtilis] SPORE GERMINATION PROTEIN GERD PRECURSOR	42 99	6.10E-41 4		1 435	424	g121132	2107-2664	5 Bt1G2636	1264 Bt1Gc1605	12
KINB SIGNALING PATHWAY ACTIVATION PROTEIN	52 99	7.20E-54 5		1. 557	557	g1708617	1967-1374	5 Bt1G2635	1264 Bt1Gc1605	12
(D64126) unknown [Bacillus subtilis]	53 100	8.00E-71 5		5 717	725	g1644216	474-1235	5 Bt1G2634	64 Bt1Gc1605	1264
(1RANSCRIPTASE BETA CHAIN) (KNA POLYMEKASE BETA SUBUNIT) [Bacillus subtilis]				•			٠			1.5
					50016	•				7
NCBI gi description	% % Ident Cvrg	-	BlastP- Prob	BlastP Score	aat_	NCBI gi	Position	Gene Id	Q Contig Id	SEQ

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	43	2.60E-24	278	226	g120294	1379-966			
99 (AJ002571) DppE [Bacillus subtilis]	35	1.70E-84	846	811	g2632017	10465-8792	3 Bt1G2669	1268 Bt1Gc1613	
100. (AF015775) YodH [Bacillus subtilis]	32	8.20E-30	330	317	g2415387	8691-7981	3 Bt1G2668	1268 Bt1Gc1613	
99 (AF015775) YodI [Bacillus subtilis]	46	7.00E-08	123	128	g2415388	7548-7934	_	1268 Bt1Gc1613	
	43	4.90E-119	1172	1036	g2619017	7997-9747			
,	40	5.40E-17	209	262	g1652288	7466-7906			
		1.20E-65	668	6/5	g2634341	6460-7257			
		9.40E-13	169	181	g1770010	6110-6365			
-	}			}·:· ·.					
100 HYPOTHETICAL 38.6 KD PROTEIN IN CWLD-GERD.	4	3.40E-79	796	715	g1723296	4199-5259	2 Bt1G2662	1267 Bt1Gc1612	
[Bacillus subtilis]			•						
99 (Z99111) molybdopterin converting factor (subunit 1)	48	1.90E-14	185	167	g2633802	3890-4123	2 Bt1G2661	1267 Bt1Gc1612	
[Bacillus subtilis]									
99 (Z99111) molybdopterin converting factor (subunit 2)	67	1.80E-48	506	532	g2633801	3427-3897	Bt1G2660	1267 Bt1Gc1612	
			•		•				
100 (Z99111) molybdopterin biosynthesis protein [Bacillus	61	1.10E-137	1348	1359	g2633799	2130-3419	Bt1G2659	1267 Bt1Gc1612	
subtilis]		1.00E-30	750	707	82033778	1022-2112		1207 511001012	
100 (700111) molyhdonterin hiosymthesis protein [Besillus	7	1 60E-05	080	080	67633708	1000-2115	B+1G2658	1267 B+1G-1617	
		;			Ç	,			
100 HYPOTHETICAL 28.5 KD PROTEIN IN SIGV-GREA	ස	1.00E-70	716	873	e3025279	295-1089	Bt1G2657	1267 Bt1Gc1612	
INTERGENIC REGION [Bacillus subtilis]									
61 HYPOTHETICAL 30.8 KD PROTEIN IN SINI-GCVT	51	3.50E-45	475	449	g1731039	6752-7236	Bt1G2656	1266 Bt1Gc1611	
INTERGENIC REGION [Bacillus subtilis]									
100 HYPOTHETICAL HELICASE IN SINI-GCVT	67	2.30E-206	1996	1976	g1731040	5083-6753	Bt1G2655	1266 Bt1Gc1611	
		-						•	
(GLYCINE CLEAVAGE SYSTEM T PROTEIN) [Bacillus			-						
100 PROBABLE AMINOMETHYLTRANSFERASE	67	1.50E-126	1243	1274	g1730258	4686-3601	Bt1G2654	1266 Bt1Gc1611	
P-PROTEIN) [Bacillus subtilis]			٠					•	
DECARBOXYLASE) (GLYCINE CLEAVAGE SYSTEM									
(DECARBOXYLATING) SUBUNIT 1 (GLYCINE				,					
100 PROBABLE GLYCINE DEHYDROGENASE	73	2.20E-164	1600	1689	g1730256	3571-2227	Bt1G2653	1266 Bt1Gc1611	
P-PROTEIN) [Bacillus subtilis]							-		
DECARBOXYLASE) (GLYCINE CLEAVAGE SYSTEM					٠			~	
(DECARBOXYLATING) SUBUNIT 2 (GLYCINE	•		-			•	-	·.	
			•	Score				2	
% NCBI gi description	% 9 Ident C	BlastP- Prob	BlastP Score		NCBI gi	Position	Gene Id	ID Contig Id	
								GFO	
		Table 1				, .	٠		

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(AB001488) SIMILAR TO BICYCLOMYCIN	100	46.42	9.20E-38 4.80E-80	403 804	401 892	g1881374	6024-7226	8 Bt1G2692	1272 Bt1Gc1618	
[Arabidopsis thaliana] (Z79580) putative orf [Bacillus subtilis]	_		4.90E-142	1389	1363	g1620930	2140-3546			• -
[Arabidopsis thaliana] (AL022603) putative NADPH quinone oxidoreductase	92	5 33	2.20E-36	392	204	g3080402	1127-1	8 Bt1G2689	1272 Bt1Gc1618	_
protein from Vigna radiata and a member of the zinc-binding dehydrogenase family PF 00107. ESTs gb T43674, gb H77006 and gb AA395179 come from this gene.	•	-								
[Synechocystis sp.] (AC005990) Strong similarity to gb U20808 auxin-induced	100	7 32	3.10E-37	400	247	g4056456	1127-219	8 Bt1G2688	1272 Bt1Gc1618	_
(D90904) sensory transduction histidine kinase	27	31	1.30E-28	330	269	g1652287	5741-5009	7 Bt1G2687	1271 Bt1Gc1617	_
(Z83337) ywpD [Bacillus subtilis]	100		5.70E-45	473	433	g1763705	4756-3920	7 Bt1G2686	1271 Bt1Gc1617	_
megaterium] HYPOTHETICAL 21.7 KD PROTEIN IN BETT-PRPR	100	5 39	2.80E-36	391	385	g2492902	3295-3897	7 Bt1G2685	1271 Bt1Gc1617	
(AF109909) 3-ketoacyl-CoA reductase PhaB [Bacillus	100	73	1.00E-77	782	911	g4160474	912-1652	7 Bt1G2684	1271 Bt1Gc1617	
(AF109909) PhaR [Bacillus megaterium]	100		1.50E-35	384	306	g4160473	267-1151	7 Bt1G2683	1271 Bt1Gc1617	_
(U67921) REP [Bacillus thuringiensis israelensis]	53	5 67	1.70E-36	393	553	g1619836	7349-8116	4 Bt1G2682	1270 Bt1Gc1614	
Mob protein - Bacillus sp. plasmid pTB19 [Plasmid pTB19]	100	32	3.40E-40	428	439	g80336	4123-5322	4 Bt1G2681	1270 Bt1Gc1614	_
(U67921) REP [Bacillus thuringiensis israelensis]	71		3.30E-88	881	963	g1619836	1-805	4 Bt1G2680	1270 Bt1Gc1614	. .
CHEMOTAXIS PROTEIN CHEA [Listeria	19	54	4.30E-24	284	258	g2500757	13200-12837	5 Bt1G2679	1269 Bt1Gc1615	· -
(Y08031) cheD [Bacillus cereus]			1.70E-77	780	833		12502-11939			
(Y08031) cheD [Bacillus cereus]		7 89	1.70E-77	780	833		12502-11939	5 Bt1G2677	1269 Bt1Gc1615	·
aeolicus										
(AE000745) flagellar switch protein FliN [Aquifex		_	0.00017	91	113		11350-10272			_
[Treponema pallidum] (M80245) CheR [Bacillus subtilis]	10,) 47	4.00E-60	616	595	g143805	8074-8850	5 Bt1G2675	1269 Bt1Gc1615	_
(figk) [Helicobacter pylori 26695] (AE001240) flagellar hook-associated protein 1 (figk)	54	23	1.90E-22	269	126	g3322960	6436-3613	5 Bt1G2674	1269 Bt1Gc1615	_
[Bacillus subtilis] (AE000618) flagellar hook-associated protein 1 (HAP1)	36	5 24	6.70E-16	208	120	g2314271	6439-4138	5 Bt1G2673	1269 Bt1Gc1615	_
FLAGELLAR HOOK-ASSÖCIATED PROTEIN 3 (HAP3)	100	5 27	1.60E-26	299	236	g2829830	5083-4221		1269 Bt1Gc1615	
(TIE100E) I for (Vibria namboumalutions)			4 10E 1	163	3	~1777365	191/ 1209	5 B+1C3671	•	.
NCBI gi description	% % Ident Cvrg	% Ident	BlastP- Prob	BlastP Score	nap Score	NCBI gi	Position	Gene Id	ID Contig Id	- _ 9
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Table 1

HYPOTHETICAL PROTEIN HI0143 [Haemophilus influenzae Rd]		27	1.80E-25	289	244	g2507342	3152-2292	Bt1Gc1624 Bt1G2712	1276 1
5'REGION (ORFB) [Clostridium perfringens]		2/	0.40E-30	331	. 00	8141080	3140-2301	B11021024 B1102/11	12/01
(Z99104) similar to hypothetical proteins [Bacillus subtilis]	8 8	ያ 4	8.40E-60	613	712	g2632437	1990-1079		
(AB002150) YbbF [Bacillus subtilis]	80	51	5.20E-67	681	854	g1256135	1075-1		
[Bacillus halodurans]								-	
(AB011836) response regulator aspartate phosphatase	98	24	5.60E-31	341	275	g4512351	8057-9110	Bt1Gc1621 Bt1G2708	1275 1
gene tnpA [Staphylococcus aureus]	100	37	1.60E-63	648	555	g224807	7670-6579	Bt1Gc1621 Bt1G2707	1275 I
butyricum]									
hypothetical protein - Clostridium butyricum [Clostridium	100	28	7.20E-70	708	478	g481912	6610-4600	Bt1Gc1621 Bt1G2706	1275 I
[Staphylococcus aureus]							• 4•		
TRANSPOSASE B (TRANSPOSON TN554)	100	29	6.30E-71	718	459	g135956	6597-4642	Bt1Gc1621 Bt1G2705	1275 I
butyricum]				•					
hypothetical protein - Clostridium butyricum [Clostridium	99	34	2.00E-10	147	156	g481913	4568-4197	Bt1Gc1621 Bt1G2704	1275 I
(AF015825) unknown [Bacillus subtilis]	100	63	7.70E-50	519	828	g2612912	2499-3260	Bt1Gc1621 Bt1G2703	1275 I
(Y11477) endolysin [Bacteriophage Bastille]	87	47	2.30E-73	741	600	g1865711	123-2195	Bt1Gc1621 Bt1G2702	1275 I
NIFS PROTEIN HOMOLOG [Bacillus subtilis]	42	50	3.10E-37	400	388	g585557	9022-9516	Bt1Gc1620 Bt1G2701	1274 I
SYNTHETASE B) [Bacillus subtilis]									`
L-ASPARTATE OXIDASE (QUINOLINATE	100	43	1.80E-114	1129	1051	g585537	8866-7021	1274 Bt1Gc1620 Bt1G2700	1274 I
STRESS PROTEIN 70) (GSP70) [Bacillus subtilis]						٠.			
(DECARBOXYLATING)) (QAPRTASE) (GENERAL									
(QUINOLINATE PHOSPHORIBOSYLTRANSFERASE								-	
PYROPHOSPHORYLASE (CARBOXYLATING)									
PROBABLE NICOTINATE-NUCLEOTIDE	100	57	5.50E-79	794	843	g3183539	7317-6448	1274 Bt1Gc1620 Bt1G2699	1274 I
(Z99118) quinolinate synthetase [Bacillus subtilis]	100	67	4.20E-129	1267	1329	g2635250	6453-5350	Bt1Gc1620 Bt1G2698	1274 I
	٠.								
	99	35	7.80E-25	283	240	g1941918	1503-1000	Bt1Gc1620 Bt1G2697	1274 I
[Bacillus subtilis]		i		.,					
(Z99108) similar to iron(III) dicitrate transport permease	100	32	5.10E-37	398	280	£2633168	12-950	Bt1Gc1620 Bt1G2696	1274 F
(X61953) abrB [Bacillus subtilis]	%	53	1.50E-21	252	218	g39805	7479-7201	Bt1Gc1619 Bt1G2695	1273 I
$\overline{}$	37	24	7.20E-09	138	100	g3005554	1-367	Bt1Gc1619 Bt1G2694	1273 I
family) [Bacillus subtilis]									
(Z99105) similar to transcriptional regulator (AraC/XylS	100	51	4.10E-74	748	778	g2632518	8717-9610	Bt1Gc1618 Bt1G2693	1272 I
RESISTANCE PROTEIN [Racillus subtilis]									
NCBI gi description	Cyrg	70 Ident	Prob	Score	nap Score	NCBI gi	Position	Contig Id Gene Id	85,
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1284 Bt1Gc1631	1284 BtlGc1631	1284 Bt1Gc1631	1284 Bt1Gc1631	1284 Bt1Gc1631		1283 Bt1Gc1627	1283 Bt1Gc1627	1283 Bt1Gc1627	• .	1282 Bt1Gc1629	1282 Bt1Gc1629			1282 Bt1Gc1629	1282 Bt1Gc1629	1282 Bt1Gc1629	1282 Bt1Gc1629	•	1282 Bt1Gc1629	1281 Bt1Gc1628	1281 Bt1Gc1628	÷		1281 Bt1Gc1628		1281 Bt1Gc1628	1281 Bt1Gc1628	1281 Bt1Gc1628	1280 Bt1Gc1623	1280 Bt1Gc1623		1280 Bt1Gc1623	SEQ ID Contig Id NO
1 Bt1G2760	1 Bt1G2759	1 Bt1G2758	1 Bt1G2757	1 Bt1G2756		7 Bt1G2755	7 Bt1G2754	7 Bt1G2753		9 Bt1G2752	9 Bt1G2751			9 Bt1G2750	9 Bt1G2749	9 Bt1G2748	9 Bt1G2747		9 Bt1G2746	8 Bt1G2745	8 Bt1G2744			8 Bt1G2743		8 Bt1G2742	8 Bt1G2741	8 Bt1G2740	3 Bt1G2739	3 Bt1G2738		3 Bt1G2737	Gene Id
6267-4331	3466-4734	3114-1348	1367-630	494-59		8514-7591	7495-6035	3015-348		7975-5858	5752-4574			4521-3661	3590-2457	2444-1308	1220-1	-	454-1	8661-8076	8025-6562			4471-5859		4344-3419	1227-2824	1-1158	7944-8318	7580-6393		6379-5540	Position
g4835822	g1074652	g1770001	g1770002	g1770003		g417116	g98699	g2337795		g1176956	g1303931	,		g1170972	g1168282	g1168281	g2983006		g133449	g3687663	g3687664	•	•	g1881332	•	g401589	g2293261	g1934815	g2293276	g2293329		g2293328	NCBI gi
632	899	1944	797	426		1045	1565	1509		2906	1252			958	1275	1587	95		360	327	819			922		445	1234	1385	414	1541		1137	nap Score
617	809	1747	827	373		1057	1538	1630		2931	1172			965	1250	1443	154		285	414	837		•	937		494	1170	1202	421	1270		1062	BlastP Score
3.20E-60	1.40E-80	5.70E-180	1.80E-82	2.30E-34		7.50E-107	8.00E-158	1.40E-167		1.90E-305	4.90E-119			4.20E-97	2.60E-127	9.30E-148	3.60E-11		4.80E-25	1.00E-38	1.50E-83			3.90E-94		3.40E-47	7.90E-119	3.20E-122	1.90E-39	2.00E-129		2.20E-107	BlastP- Prob
33	4	65	66	62		61	66	40		77	61			2	66	80	30		50	42	38			43		33	62	67	70	75		76	% Ident
95	. 100	100	100	99	. :	100	100	100		100	100			100	100	100	61	-	89	85	100		_	100			100	86	99	_		100	% Cvrg
	hypothetical protein HI1104 - Haemophilus influenzae (strain Rd KW20) [Haemophilus influenzae Rd]	(Z75208) autolysin sensor kinase [Bacillus subtilis]	(Z75208) autolysin response regulator [Bacillus subtilis]	(Z75208) hypothetical protein [Bacillus subtilis]	(HEME SYNTHETASE) [Bacillus subtilis]	FERROCHELATASE (PROTOHEME FERRO-LYASE)	catalase (EC 1.11.1.6) - Listeria seeligeri []	(Y13937) putative PacL protein [Bacillus subtilis]	5'REGION [Bacillus subtilis]	HYPOTHETICAL 79.2 KD PROTEIN IN ACDA	(D84432) YqiL [Bacillus subtilis]	DEHYDROGENASE) (BHBD) [Bacillus subtilis]	DEHYDROGENASE (BETA-HYDROXYBUTYRYL-COA	PROBABLE 3-HYDROXYBUTYRYL-COA	ACYL-COA DEHYDROGENASE [Bacillus subtilis]	ACYL-COA DEHYDROGENASE [Bacillus subtilis]	(AE000683) putative protein [Aquifex aeolicus]	SUBUNIT [Bacillus subtilis]	DNA-DIRECTED RNA POLYMERASE DELTA	(AF049873) response protein [Lactococcus lactis]	(AF049873) sensor protein [Lactococcus lactis]	MELILOTI. [Bacillus subtilis]	CATABOLISM (MOCR) GENE OF RHIZOBIUM	(AB001488) SIMILAR TO THE RHIZOPINE	INTERGENIC REGION [Escherichia coli]	HYPOTHETICAL 33.1 KD PROTEIN IN SELC-NLPA	(AF008220) YtsJ [Bacillus subtilis]	(Z93937) unknown [Bacillus subtilis]	(AF008220) YtcD [Bacillus subtilis]	(AF008220) YtbD [Bacillus subtilis]	subtilis]	(AF008220) putative morphine dehydrogenase [Bacillus	NCBI gi description

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SEQ ID Contig Id Gene Id NO	Position	NCBI gi	nap Score	BlastP Score	BlastP- Prob	% Ident	Cvrg	NCBI gi description
1285 Bt1Gc1630 Bt1G2761	79-1322	g732364	807	735	9.90E-73	43	91	HYPOTHETICAL 50.0 KD PROTEIN IN UNG-ROCA
1285 Bt1Gc1630 Bt1G2762	1591-2394	g401595	456	412	1.70E-38	. 37	100	HYPOTHETICAL 29.7 KD PROTEIN IN IBPA-GYRB
1285 Bt1Gc1630 Bt1G2763	3634-2960	g731058	770	770	1.90E-76	2	100	INTERGENIC REGION [Escherichia coli] URACIL-DNA GLYCOSYLASE (UDG) [Bacillus subtilis]
1285 Bt1Gc1630 Bt1G2764		g1724002	701	743	1.40E-73	47	100	HYPOTHETICAL ABC TRANSPORTER ATP-BINDING PROTEIN 2 IN GLPD-CSPB INTERGENIC REGION
								[Bacillus subtilis]
1285 Bt1Gc1630 Bt1G2765	6322-5568	g1730938	278	348	1.00E-31	31	100	HYPOTHETICAL 28.9 KD PROTEIN IN ILVA 3'REGION
1285 Bt1Gc1630 Bt1G2766	7422-6448	g2633758	643	637	2.40E-62	40	100	[Bacillus subtilis] (Z99111) similar to transcriptional regulator (LacI family)
1285 Bt1Gc1630 Bt1G2767	9471-8177	g118592	1513	1452	1.00E-148	68	100	HOMOSERINE DEHYDROGENASE (HDH) [Bacillus subtilis]
1286 Bt1Gc1632 Bt1G2768	497-323	g2634358	151	162	5.20E-12	52	98	(Z99114) yozD [Bacillus subtilis]
Bt1Gc1632	1466-816	g2415400	392	410	2.70E-38	38	100	(AF015775) YodN [Bacillus subtilis]
1286 BHGC1632 BHG2771	7301-2008	876054500	310	270	1.90E-14	2 6	8.4	(ABOOGGGS) Volti [Bacillus subtilis]
Bt1Gc1632	3698-2286	g4033499	2025	1949	2.20E-201	77	100	HYPOTHETICAL 54.1 KD PROTEIN IN DEOD-ARGE
								INTERGENIC REGION [Bacillus subtilis]
Bt1Gc1632	5294-3909	g1064808	843	870	4.90E-87	42	100	(D78193) positive regulatory protein [Bacillus subtilis]
Bt1Gc1632	6584-5522	g2415402	434	472	7.30E-45	39	100	(AF015775) YodP [Bacillus subtilis]
1286 Bt1Gc1632 Bt1G2776	10045-7630	g2529465	1371 7 57	1308	1.90E-133	3 8	3 5	(AF006665) YokP [Bacillus subtilis]
pt10c107	10040-7000	E400/01/	727	000	J.00E-00	3	5	ketoacid-CoA transferase precursor [Homo sapiens]
1286 Bt1Gc1632 Bt1G2777	10291-8990	g2529462	1096	1076	7.20E-109	51	200	(AF00665) YokM [Bacillus subtilis]
		q					ò	[Bacillus thuringiensis]
1287 Bt1Gc1633 Bt1G2779	659-99	g1/31021	430	352	3.80E-32	45	99	HYPOTHETICAL 21.4 KD PROTEIN IN SODA-COMGA INTERGENIC REGION [Bacillus subtilis]
1287 Bt1Gc1633 Bt1G2780	3346-1570	g1653786	880	942	1.10E-94	32	71	(D90916) mannose-1-phosphate guanyltransferase
1287 Bt1Gc1633 Bt1G2781	4334-3667	g2983430	326	286	3.80E-25	37	95	[Synechocystis sp.] (AE000713) transcriptional regulator (PhoU-like) [Aquifex
1287 Bt1Gc1633 Bt1G2782	5393-4638	g3024474	887	890	3.70E-89	65	100	aeolicus] PROBABLE PHOSPHATE TRANSPORT ATP-BINDING
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*	SEQ ID Contig Id Gene Id NO	Position	NCBI gi	nap Score	BlastP Score	BlastP- Prob	%: Ident C	%Cvrg	NCBI gi description
	1290 Bt1Gc1637 Bt1G2804	2545-2198	g586872	378	411	2.10E-38	60	99	HYPOTHETICAL 14.1 KD PROTEIN IN XPAC-ABRB
	1290 Bt1Gc1637 Bt1G2805	3387-2563	g586871	1147	1147	2.20E-116	76	100	INTERGENIC REGION [Bacillus subtilis] HYPOTHETICAL 31.2 KD PROTEIN IN XPAC-ABRB
	1290 Bt1Gc1637 Bt1G2806	4376-3396	g586870	862	816	2.60E-81	53	100	INTERGENIC REGION [Bacillus subtilis] DNA POLYMERASE III, DELTA' SUBUNIT [Bacillus
	1290 Bt1Gc1637 Bt1G2807	5038-4403	g586867	579	591	1.80E-57	52-	100	SUBSTITUTE
	1290 Bt1Gc1637 Bt1G2808	6452-4943	g586866	1006	1079	3.50E-109	49	100	HYPOTHETICAL 53.2 KD PROTEIN IN XPAC-ABRB
	1291 Bt1Gc1638 Bt1G2809	1-863	g586901	1183	1201	4.10E-122	81	63	DNA REPAIR PROTEIN RADA HOMOLOG (DNA REPAIR PROTEIN SMS HOMOLOG) [Bacillus subtilis]
	1291 Bt1Gc1638 Bt1G2810	870-1940	g586902	1136	1087	4.90E-110	63	100	HYPOTHETICAL 40.7 KD PROTEIN IN MECB-GLTX INTERGENIC REGION (Bacillus subtilis)
	1291 Bt1Gc1638 Bt1G2811	2104-3210	g586903	1252	1243	1.50E-126	70	100	HYPOTHETICAL 40.9 KD PROTEIN IN MECB-GLTX INTERGENIC REGION [Bacillus subtilis]
	Bt1Gc1638	3227-3919	g2127058	579	603	9.60E-59	51	100	hypothetical protein - Bacillus subtilis [Bacillus subtilis]
	1291 BUICE1838 BUICE813	0000	80470//	Q.	94/	2.10E-03	6	. 4	INTERGENIC REGION [Bacillus subtilis]
	1291 Bt1Gc1638 Bt1G2814	4593-6041	g135106	1927	1898	5.70E-196	74	100	GLUTAMYL-TRNA SYNTHETASE (GLUTAMATE-TRNA LIGASE) (GLURS) [Bacillus subtilis]
	Bt1Gc1638	7141-7984	g549024	1258	1260	2.30E-128	8 2	. 8	CYSTEINYL-TRNA SYNTHETASE (CYSTEINETRNA LIGASE) (CYSRS) [Bacillus subtilis]
	1292 Bt1Gc1634 Bt1G2817	622-1 1432-1048	g1/630/9 g2623262	603	628	9.90E-41 2.20E-61	98 8	99	(AF030979) phosphonoacetaldehyde hydrolase [Bacillus
	1292 Bt1Gc1634 Bt1G2818	4517-1454	g1469285	356	448	6.10E-42	25	80	cereus] (U05042) afuB gene product [Actinobacillus
	1292 Bt1Gc1634 Bt1G2819	3178-1475	g3319739	193	288	1.00E-22	22	100	pleuropneumoniae] (AL031035) ABC transporter integral membrane protein
	1292 Bt1Gc1634 Bt1G2820	4231-2116	g3341854	628	596	5.30E-58	4	71	[Streptomyces coelicolor] (AF077856) PotA [Actinobacillus actinomycetemcomitans]
	1292 Bt1Gc1634 Bt1G2821	5274-4249	g1469286	361	415	8.00E-39	30	00	(U05042) afuA gene product [Actinobacillus
	1292 Bt1Gc1634 Bt1G2822	7566-6217	g2633721	1396	1046	1.10E-105	60	100	pleuropneumoniae] (Z99111) similar to Na+-transporting ATP synthase

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(X98606) potential coding region [Clostridium difficile]	44 100	1.10E-112	1112	. 1137	g1418413	1225-3058	Bt1G2847	b Bt1Gc1641	1296
[Bacillus									3
(Z99112)	71 15	3.50E-60	626	586	g2633966	82-4639	Bt1G2846	6 Bt1Gc1641	1296
(Z82044) hypothetical 40.7 kd protein [Bacillus subtilis]	50 90	4.90E-71	719	811) g1673393	11135-10170	Bt1G2845	5 Bt1Gc1640	1295
embriliel		i		,	0.000				į
(782044)	_	3 20E-177	_	1905	ø1673392	8352-10116	Bt1G2844		1295
(Z82044)	74 99	1.30E-38	413	398	g1673391	7792-8107	Bt1G2843	5 Bt1Gc1640	1295
(Z99108) yfhS [Bacillus subtilis]	62 99	1.30E-24	281	281	g2633187	6663-6442	Bt1G2842	5 Bt1Gc1640	1295
(299108)	58 100	2.30E-112	1109	1142	g2633186	5300-6409	Bt1G2841	Bt1Gc1640	1295
(Z99108)		1.90E-92	_	1006	g2633185	5156-4179	Bt1G2840		1295
(Z99108)		3.00E-23	-	246	g2633177	3038-3349	Bt1G2839		1295
(Z99108) yfhG [Bacillus subtilis]	40 100	3.90E-46		508	g2633176	2209-3012	Bt1G2838	5 Bt1Gc1640	1295
(Z99108) similar to cell-division inhibitor [Bacillus subtilis]	48 100	3.90E-78	786	741	g2633175	2127-1219	Bt1G2837	5 Bt1Gc1640	1295
horikoshii]					•				
	35 97	4.60E-52	540	464	g3257222	1079-1	Bt1G2836	5 Bt1Gc1640	1295
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		9.70E-66		628		11646-12259	Bt1G2835		1294
	27 100	4.70E-34	370	314	g4481748	10424-11470	Bt1G2834	4 Bt1Gc1639	1294
PROTEIN (ORF11) [Bacillus subtilis]					ı				
	65 100	1.10E-75	763	750	g1175624	9842-10516	Bt1G2833	4 Bt1Gc1639	1294
				-					
HYPOTH	100	4.60E-114		1318	g732334	8061-9242	Bt1G2832		1294
(D86417)	45 100	2.10E-54	562	576	g2443249	5784-6581	Bt1G2831	4 Bt1Gc1639	1294
_	33 100	2.40E-46	486	507	g2443247	4796-5794	Bt1G2830	4 Bt1Gc1639	1294
HEMA INTERGENIC REGION (ORFX) [Bacillus subtilis]					٠.			-	
HYPOTHETICAL GTP-BINDING PROTEIN IN LONA-	77 49	6.50E-37	397	378	g586754	9469-9757	Bt1G2829	3 Bt1Gc1635	1293
subtilis] ATP-DEPENDENT PROTEASE LA 1 [Bacillus subtilis]	73 100	1.40E-277	2668	2995	g585415	7148-9469	Bt1G2828	3 Bt1Gc1635	1293
-	72 100	5.10E-211	2040	2089	g1708857	5289-6942	Bt1G2827	3 Bt1Gc1635	1293
(X95306)	8 7 100	6.10E-183		1835	g1296452	3923-5179	Bt1G2826	Bt1Gc1635	1293
(VEG2) [Bacillus subtilis]			٠		(
		1.90E-140	<u></u>	1577	£2829689	2381-3652	Bt1G2825		1293
(Z99121)		2.60E-08		124	£2635899	105-1	Bt1G2824		1293
(AF109909) PHA synthase PhaC [Bacillus megaterium]	70 100	1.90E-140	1374	1419	g4160475	9964-8879	Bt1G2823	2 Bt1Gc1634	1292
NCBI gi description	=		Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	8
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SEQ	1296 E		1296 E	1297 E	1297 E		1297 E		1207 1		•	1297 E	٠			1297 E		129/ 6		1298 1		1298 I	1298 I		1298 1	1298 I
Contig Id	Bt1Gc1641		Bt1Gc1641	Bt1Gc1642	Bt1Gc1642		Bt1Gc1642	*10°164	B+1G-1643	Bt1Gc1642		1297 Bt1Gc1642		,		Bt1Gc1642	· ·	Bt1Gc1642		Bt1Gc1644		Bt1Gc1644	Bt1Gc1644	, ,	Bt1Gc1644	Bt1Gc1644
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Gene Id	Bt1G2848		Bt1G2849	Bt1G2850	Bt1G2851		Bt1G2852	Coes	B+102854	Bt1G2855		Bt1G2856 11194-10394 g3121841	٠			Bt1G2857		8C872)13B		Bt1G2839		Bt1G2861	Bt1G2862		Bt1G2863	B+1 C2864
Position	3174-4640		4131-5699	893-1	1381-914		6019-1721	70//	0716-7057	10370-9211		11194-				11985-11206		12622-12068		3557-1163 9586876		431-3640	8116-6789		9274-8078	
tion	4640		5699					6163	7057	<u>8</u>		10394				11206		89071		1163		3640	6789		8078	
NCBI gi	g4322642	· : ``.	g2577963	g2634032	g418459		g118793	2624	87634078	93913544		g3121			•	g3183468		g3122/62		20340Z	(g3256896	£2661710	5	g3581864	20106777
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BlastP Score	562	-	1216	1148	435		5507	2000	1264	1276	•	681				796		633	,) 284		309	711		673	
BlastP- Prob	2.10		1.10E-123	1.70E-116	6.10			301	9.70E-210	4.60E-130		5.20				3.40	• •	6.40		9 50		1.30	3.50		3.70	4 00E-305
•	2.10E-54		-123	-116	6.10E-41		0	3	130	130		5.20E-67				3.40E-79	:	6.40E-62	} }	1.80E-30		1.30E-26	3.50E-70		3.70E-66	2
% % Ident Cvrg	37		45	8 1	69		75		\$ 6	<u></u>		55				65		75	; !	ಚ ୧		27	35		36	వ
Cvrg	88		100	80	99		100		3 5			100				100		, 99	i	3 £		39	00		100	3
	(AF093246) NAD(P)H-dependent 2-cyclohexen-1-one	reductase Ncr [Pseudomonas syringae pv. glycinea]	(Y15254) YerD protein [Bacillus subtilis]	(Z99112) nus	HYPOTHETICAL 17.6 KD PROTEIN IN NUSA	5'REGION (P15A) (ORF1) [Bacillus subtilis]	DNA POLYMERASE III, ALPHA CHAIN [Bacillus	subtilis]	(299112) pro	1-DEOXY-D-XYLIII OSE 5-PHOSPHATE	REDUCTOISOMERASE (DXP REDUCTOISOMERASE)	[Bacillus subtilis] PHOSPHATIDATE CYTIDYLYLTRANSFERASE (CDP-	DIGLYCERIDE SYNTHETASE) (CDP-DIGLYCERIDE BYRCERIDE SYNTHETASE) (CDP-DIGLYCERIDE	SYNTHASE) (CDS) (CTP:PHOSPHATIDATE	CYTIDYLYLTRANSFERASE) (CDP-DAG SYNTHASE) [Bacillus subtilis]	HYPOTHET	INTERGENIC REGION [Bacillus subtilis]	RELEASING FACTOR) (RRF) (VEGETATIVE PROTEIN	12B) (VEG12B) [Bacillus subtilis]	HYPOTHETICAL 47.7 KD PROTEIN IN METS-KSGA	INTERGENIC REGION [Bacillus subtilis]	(AP000002) 739aa long hypothetical chemotaxis protein	(AL009204) FAD-dependent oxidoreductase [Streptomyces	coelicolor]	(AL031541)	phospholipase C (EC 3.1.4.3) - Bacillus cereus (strain IAM
	NAD(I	r [Psei	rD pro	2) nusA [Bacillus subtilis]	ICAL	P15A)	MERA		2) rimilar to hypothetical proteins (Bacillus suomis)	XY-D-XYLIII OSE 5-PHOSPHATE	SOME	tilis]	DE SY) (CDS	LTRAI tilisi	THETICAL 29.8 KD PROTEIN IN FRR-CDSA	CREC	FAC	2B) [E		CREC	739aa horibo	FAD-c	,	hypoth	e C(E
NCBI gi description	P)H-de	omopr	tein []	cillus	17.6 K	(ORF)	SE III,	1	NA SY		RASE	CYT	HIN)(CTI	NSFEF	29.8 K	NOIS	TOR)	Bacillu	Kinase 47 7 K	NOIE	long h	lepend		etical	C3.1.
gi des	pende	nas sy	Bacillu	subtili	D PR) [Ba	ALPI		hatical	E S-PF	(DXP	IDYL	ETASI	P:PHO	ASE)	D PR	Bacil	GFAC	s subti		[Bacil	ypothe	ent ox		protei	1 2 1
criptio	nt 2-cy	ringae	ıs subt	ഭ	OTEIN	cillus	IA CE	; 3		HOSPI Property	REDU	YLTR		SPHA	(CDP	OTEIN	lus sut	(VEG	lis]		lus sut	etical c	idored		n SCI3	Bacill _i
ğ	/clohe	pv. gl	ilis]		NN	subtilis	AIN			ATE	JCTO	ANSF	P-DIC	TIDA	-DAG	IN F	otilis]	ETAT) - 		tilis]	hemot	uctase		5.31	
	xen-1-	ycines			USA	۳	[Bacil	L .: 1:		20114	MOSI	ERAS		TE	NAS	RR-CI		TVE P		ETS		axis p	Stre		Strep	ne (etr
	one	<u>ت</u>		_			lus	<u>.</u>	2 K	94041	ERASI	E (CD)	ERIDE		THASI	SA	•	ROTE		KSG A		rotein	ptomy		1541) hypothetical protein SCI35.31 [Streptomyces	;; I A]
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	Bt1G2886	Bt1G2885	Bt1G2884	Bt1G2883	Bt1G2882	;	Bt1G2881	Bt1G2880	Bt1G2879		Bt1G2878	Bt1G2877	Bt1G2876	Bt1G2875	Bt1G2874	Bt1G2873	Bt1G2872	Bt1G2871	Bt1G2870		Bt1G2869	Bt1G2868	Bt1G2867		Bt1G2866	Bt1G2865	Gene Id
	2886	2885		2883	2882		2881	2880	_		2878	2877	2876	2875	2874	2873	2872	2871	2870		2869	2868	2867		2866		e Id
	6820	Ξ	12149	9339-9731	8284		7096-7623	. 190	10177		9002	8164-8931	25-	7668-6526	6170-5061	4995-3085	2990-528	7734-5337	5561-4850		2157-3320	1519.	869-	_	692-1	11626	Position
	6820-1254	1-353	12149-11225	-9731	8284-9207	į	7623	1902-1	10177-10627		9002-9932	-8931	25-579	-6526	5061	-3085	-528	-5337	4850		3320	1519-2160	869-1498		2-1	11626-11911	tion
	g2622173	g115692	g115950	g1763704	g984656	q	o 1788724	g3036999	-		g732393	g729326	g1001205	g118797	g132246	g121887	g121881	g4104606	g4481749		g728801	g728800	g728799		g728788	g420808	Z.
	2173	692	950	3704	656		8724	6999	1269		393	326	1205	797	246	887	88 1	4606	1749		801 1	800	799		788	808	NCBI gi
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	1.40E-55	4.80E-25	4.70E-121	6.80E-16	4.00E-69		1 40F-16	2.50E-37	7.60E-27		1.10E-87	6.60E-28	6.60E-28	3.20E-147	6.90E-120	1.30E-258		1.30E-40	1.40E-50		3.10E-133	1.30E-52	1.60E-74		3.70E-82	7.50E-20	BlastP- Prob
	55	.25	21	<u>.</u> 16	69	,	<u>,</u>	37	27		87	28	28	47	20	58	0	4	50		33	52	74		82	20	
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	Plasmid AE0008	(CATABOLITE CONTROL PROTEIN) [Bacillus subtilis] CHLORAMPHENICOL ACETYLTRANSFERASE	mechanosensitive channel protein [Bacillus subtilis] GLUCOSE-RESISTANCE AMYLASE REGULATOR	(Z83337)	(D25292) ORF3 [Salmonella typhimurium]	Escherichia coli]	(AF000326) nutative 2-component transcriptional regulator	(AF019629) putative fimbria-associated protein	(AB001488) FUNCTION UNKNOWN. [Bacillus subtilis]	INTERGENIC REGION (ORF.1) [Bacillus subtilis]	[Bacillus megaterium] HYPOTHETICAL 34.3 KD PROTEIN IN BGLH-WAPA	GLUCOSE 1-DEHYDROGENASE III (GLCDH-III)	(D64003	NA F	RECF PROTEIN [Bacillus subtilis]	DNA GYRASE SUBUNIT B [Bacillus subtilis]	DNA GYRASE SUBUNIT A [Bacillus subtilis]	(AF036967) putative histidine kinase [Lactobacillus sakei]	(AF007865) BacR [Bacillus licheniformis]	subtilis]	subtilis] ACETO	subtilis] ACETO	ACETOIN UTILIZATION ACUA PROTEIN [Bacillus	COA LIGASE) (ACYL-ACTIVATING ENZYME) (ACETYL-COA SYNTHASE) [Bacillus subtilis]	aquaticus ACETYL-COENZYME A SYNTHETASE (ACETATE-	hypothetical 12K protein - Thermus aquaticus [Thermus	
	id NR79] 0878) put	RAMI	nosens OSE-I	87) hig	92) OF	richia	omyce 0326)	9629)	1488)	GENI	us meg	OSE 1)3) hy	IX TO	PROT	3YRA	3YRA	5967)	7865)				O NIC	YL-C	YL-CC	etical	
	79] putati	PHEN	itive o	highly similar to E. coli large conductance	E3 [3	coli]	s naes	putati	FUNC	CRE	gateriu ICAL	-DEH) hypothetical protein [Synechocystis sp.]	VERA	EZ	SE SU	SE SU	putati	BacR		N UTILIZATION ACUC PROTEIN [Bacillus 🗧	N UTILIZATION ACUB PROTEIN [Bacillus	TILIZ	E) (A)	DENZ	12K p	•
	ve me	ICOL	hanne TANC	milar 1	Salmo	i	lundii	ve fim	TION	GION	Im] 34.3]	YDR	ical pi	SE II	Bacill	NAR	NOBL	ve hist	[Baci		ATIC	'ATIC	ATIC	ANTH CXL-7	YME	rotein	NCB
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	tein []	RANS	Sacillu SE RU	ge coi	mirun		anscr.	ited pr	N.	acillu	N N	III (G	echoc	MA		lus su	lus su	Lac	ormis		ROTE	ROTE	ROTE	illus s	TASI	aquati	tion
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	[Plasmid NR79] (AE000878) putative membrane protein [Methanobacterium	otilis]					lator		tilis]		PA			DNA POLYMERASE III, BETA CHAIN [Bacillus subtilis]			خو	ikei]		٠.	5 √:	S	SI		TH.	. SI	· .
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Contig Id		Bt1Gc1647	Bt1Gc1648	Bt1Gc1648	Bt1Gc1648	Bt1Gc1648	! !	Bt1Gc1648	Bt1Gc1648		Bt1Gc1648	940130119	Bt1Gc1648	ें Bt1Gc1651	Bt1Gc1651	Bt1Gc1651		Bt1Gc1651	Bt1Gc1651	Bt1Gc1651	Bt1Gc1651	Bt1Gc1651	Bt1Gc1651	Bt1Gc1651	Bt1Gc1653	Rt1Gc1657	֚֚֚֓֡֜֝֜֜֝֓֜֝ ֓
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Gene Id		Bt1G2887	Bt1G2888	Bt1G2889	Bt1G2890	Bt1G2891)	Bt1G2892	Bt1G2893	!	Bt1G2894	66970119	Bt1G2896	Bt1G2897	Bt1G2898	Bt1G2899		Bt1G2900	Bt1G2901	Bt1G2902	Bt1G2903	Bt1G2904	Bt1G2905	Bt1G2906	Bt1G2907	į	R0675119
bId		2887	888	2889	2890	2891		2892	2893		2894	.093		2897	2898	2899		2900	1062	2902	2903	2904	2905				2908
Position		7991-8743	145-643	610-1648	1978-3531	5064-6536		6865-8397	6847-8144		8532-9587	0//3-10400	8337-11407	1201-1	1404-	2340-2897	•	4647-4238	5069-4830	6860-5772	7959-6880	10964	10452-9646	12176-	10231-	<u>,</u>	999-1
tion		8743	23	648	3531	6536		8397	8144		9587	0400	11407	<u>-</u>	404-2267	2897		4238	4830	5772	6880	10964-7959.	-9646	.12176-11080	10231-12365		<u> </u>
NC	٠,	g2635679	g2226153	g2226152	g2808436	g2326750		g267023	g586884		g1652052	8121030	g4218533	g2293313	g3123307	g1731066		g2635860	g2226143	g3290176	g3290177	g3290175	g2098612	g1177016	g3150046	٠.	2586848
NCBI gi		5679	6153	6152	8436	6750		023	884		2052	000	8533	3313	3307	1066		5860	6143	0176	0177	0175	8612	7016	0046		848
aat_ nap Score		457	362	1394	642	1465	!	281	416)	245	113	274	1329	974	218		385	222	775	356	860	342	1258	2253		1383
BlastP Score		:		•						•		-															
stP		496	387	1355	783	988		372	481	,	353	197	410	958	1002	291		411	225	691	361	730	433	1111	2252		1339
BlastP. Prob		2.10E-47	7.40E-36	2.00E-138	8.10E-78	1.50		2.90E-34	8.10E-46))	3.00E-32	2.00	8.30	2.30E-96	5.00E-101	1.10		2.10	1.10	4.50	4.20	3.30	9.90E-41	1.40E-112	1.70E-233		9.80E-137
		E-47	E-36	-138	E-78	1.50E-99		E-34	E-46	}	E-32	2.00E-14	8.30E-38	E-96	-101	1.10E-25		2.10E-38	1.10E-18	4.50E-68	4.20E-33	3.30E-72	E-41	-112	-233		-137
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	therr	(Z99	1 1 1	(Y1408)	Ω.	(Z98	tube	STAGE	HYT	I N	(D9)	GLU	(AJO	(AFC	TYH	H H	INI	(Z99 [Bac	(Y1407)	(AF067 cereus]	(AF067 cereus]	(AF067	(J)	IAH IAH	(AF	subs	HYH
	thermoautotrophicum]	(Z99120) similar to sepiapterin reductase [Bacillus subtilis]	(Y14080) hypothetical protein [Bacillus subtilis]	080) (080)	(AL021309) fadD13 [Mycobacterium tuberculosis]	(Z98268) hypothetical protein Rv1707 [Mycobacterium			HYPOTHETICAL 57.4 KD PROTEIN IN MFD-DIVIC	INTERGENIC REGION [Bacillus subtilis]	(D90902) hypothetical protein [Synechocystis sp.]	ENDOGLOCANASE FRECORSOR (ENDO-1,4-BETA- GLUCANASE) (ALKALINE CELLULASE) [Bacillus sp.]	(AJ010312) endo-beta-N-acetylglucosaminidase	[Streptococcus pneumoniae] (AF008220) YtiP [Bacillus subtilis]	HYPOTHETICAL OXIDOREDUCTASE IN APRE-COMK	INTERGENIC REGION (ORFX) [Bacillus subtilis] HYPOTHETICAL 19.7 KD PROTEIN IN GLNO-ANSR	INTERGENIC REGION [Bacillus subtilis]	(Z99121) similar to hypothetical proteins from B. subtilis [Bacillus subtilis]	1079)				(U66614) putative transposase [Marinococcus halophilus]	HYPOTHETICAL 43.7 KD PROTEIN IN KATB	(AF016634) ClpB chaperone homolog [Lactococcus lactis.	subsp. cremoris]	HYPOTHETICAL 40.1 KD GTP-BINDING PROTEIN IN
	otropi	simile	hypor	0) hypothetical protein [Bacillus subtilis]	9) fac	hypot	is]	SPO	ETIC	NIC	hypo	ASE)	2) enc	O Yt	ETIC	ETIC	NIC	simil: subtili	9) hypothetical protein [Bacillus subtilis]	645) spore germination protein GerIB [Bacillus	645) spore germination protein GerlC	645) spore germination protein GerIA [Bacillus	putat	ETIC	<u>4</u> ان (4	moris	ETIC
7	hicun	er to s	thetic	thetic	1013	hetic		RUL	AL 5	REG	thetic	(AL	lo-be	pneu ¡P [E	AL C	REG AL 1	REG	is] ar to l	thetic	ore g	ore g	ore g	ive tr	AL 4	pB cl	<u></u>	AĽ4
NCBI gi description	으	epiaj	al pr	al pr	3	al pro		ATIC	7.4 K	Š	al pr	KAL	ta-N-	moni: Bacill	ЫX	9.7 K	<u>N</u>	hypot	al pr	ermir	ermir	ermir	anspo	3.7 K	naper).
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		subt			ت	erium		V SPORULATION PROTEIN B [Bacillus subtilis]	ÍVIC			illus			E-CO	s] SNSI		ubtili		cillus	[Bacillus	cillus	yphilı		us lac		Z
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HYPOTHETICAL 40.7 KD PROTEIN IN CSPB-GLPP INTERGENIC REGION [Bacillus subtilis]	100	1 31	3.40E-31	343	391	g1724005	10838-9774 g1724005	Bt1Gc1654 Bt1G2927	1309 Bt1Gc1	
PHOSPHATE KINASE) (HMP-P KINASE) [Bacillus subtilis]							•			
PHOSPHOMETHYLPYRIMIDINE KINASE (HMP-	100	1 66	2.80E-91	910	896	g732357	8869-9678	654 Bt1G2926	1309 Bt1Gc1654	
(Y14079) hypothetical protein [Bacillus subtilis]	100	5 48	3.10E-85	853	778	g2226140	8731-7142	654 Bt1G2925	1309 Bt1Gc1654	
[Strentomyces coefficials]	٠		i			0				
(AI 049754) purative two-component system regulator	<u>.</u>	8 44	6 20F-48	501	460	p4753864	7168-6483	654 Bt1G2924	1309 Bt1Gc1654	
(A1027606) acyt cantet protein priospriodicsterase (Bacillas			1.5015-07		è	64017074				
(AF007868) and carrier protein phosphodiasterase [Bacillus	.	7	1 505-6	787	766	02619052	6303-5680	654 Rt1G2923	1309 Rt1Gc1654	
HYPOTHETICAL 25.3 KD PROTEIN IN NEO-FROA	100	0 42	1.60E-40	431	410	g463600	4983-3042	034 BIIU2922	F309 BUIGCI034	
(D86418) YmB [Bacillus subtilis]	97		6.30E-71		. 794	g2116/60	4836-4145			
meyeri]	.			!						
(Y10744) O-acetylhomoserine sulfhydrylase [Leptospira	100	9 . 48	7.20E-109	1076	1006	g1835113	2374-1073	654 Bt1G2920	1309 Bt1Gc1654	
coli]						•				
(HOMOSERINE O-TRANSSUCCINYLASE) [Escherichia										
HOMOSERINE O-SUCCINYLTRANSFERASE	100	3 50	5.20E-83	832	812	g127029	1065-139	654 Bt1G2919	1309 Bt1Gc1654	
[Borrelia burgdorferi]									•	
FLAGELLAR MOTOR SWITCH PROTEIN FLIG	100	7 35	4.40E-47	493	459	g1706844	12123-11098	652 Bt1G2918	1308 Bt1Gc1652	
(AEUU1218) Ilagellum-specific A1P synthase (IIII)	100	40	9.00E-79	/92	728	g3322083	8806-77401	027 8(107817	1208 BITGG1027	
(A0/136) Hagelilli [Dacillus muringlensis]	. 0		2001-200		7099	81004095	10/52 0000			
PROTEIN THDF [Bacillus subtilis]	, . 0		1 005 10		1000	~192/20¢	1 716			
POSSIBLE THIOPHENE AND FURAN OXIDATION	001	5 73	8.10E-165	1604	1709	g135725	9912-8545	657 Bt1G2914	1307 Bt1Gc1657	
subtilis]				÷					,	
SUBTRIES SUBTRIES STATES A STATE OF SUBTRIES SUB	100	1 82	2.80E-281	2703	2703	g1211 87	8495-6612	657 Bt1G2913	1307 Bt1Gc1657	
GLUCOSE INHIBITED DIVISION PROTEIN B [Bacillus	100	9 70	3.40E-79	796	887	g121190	6584-5868	657 Bt1G2912	1307 Bt1Gc1657	
INTERGENIC REGION [Bacillus subtilis]			٠							
HYPOTHETICAL 32.8 KD PROTEIN IN SPO0J-GIDB	100	7 67	1.60E-97	969	940	g586854	5759-4896	657 Bt1G2911	1307 Bt1Gc1657	
SOJ PROTEIN [Bacillus subtilis]	100	7 81	2.80E-107	1061	1061	g586852	4696-3938	657 Bt1G2910	1307 Bt1Gc1657	
INTERGENIC REGION [Bacillus subtilis]					٠.	٠				
HYPOTHETICAL 22.5 KD PROTEIN IN RPSF-SPO0J	100	8 48	6.20E-48	501	484	g586851	2474-3088	657 Bt1G2909	1307 Bt1Gc1657	
RPSF-SPO0J INTERGENÍC REGION [Bacillus subtilis]									2	
	CALS	Ident	Frob	Score	Score				_	
NCBI gi description	8	I %	BlastP-	BlastP		NCBI gi	Position	Id Gene Id	D Contig Id	
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SENSOR KINASE CITA [Klebsiella pneumoniae] METHYL-ACCEPTING CHEMOTAXIS PROTEIN MCPA			4.70E-82 7.20E-125	—	756 1440	g1705888 g730002	3813-1427 5171-3189			· :
	100	41	6 80F-42	444	44	02443739	1422-742	Bt1G2949	1313 Br1Gc1659	-
	5 54	71 65	1.10E-71	725	765	g1177013	1-691	Bt1G2948	1313 Bt1Gc1659	<u>,</u>
	86 98	-	1.90E-204	1978	2239	g118589	12077-10552	Bt1G2947	1312 Bt1Gc1660	<u> </u>
	50 100		1.70E-100	997	987	g1323764	8584-9726	Bt1G2946	1312 Bt1Gc1660	
	8 100	66 38	1.10E-66	678	841	g1731094	8299-6903	Bt1G2945	1312 Bt1Gc1660	
(AJ002571) YkfB [Bacillus subtilis]	5 100	-	3.10E-53		545	g2632019	5522-4434			<u> </u>
	oo I oo	22 23	1./0E-22	201	122	g323/120	2/93-3363	Bt102943	1317 RITAC1000	<u>.</u>
				•	<u>;</u>					•
	2 99	30 42	1.50E-30	337	245	g465586	2187-2984	Bt1G2942	1312 Bt1Gc1660	=
	9 100	33 79	6.40E-133	1303	1354	g2633127	841-1815	Bt1G2941	1312 Bt1Gc1660	=
	1 17	08 41	4.30E-08		107	g224807	178-1	Bt1G2940	1312 Bt1Gc1660	=
INTERGENIC REGION [Bacillus subtilis]						•				
HYPOTHETICAL 45.5 KD PROTEIN IN BGLS-KATB	0 38	53 70	1.90E-53	553	604	g1177013	13146-13639	Bt1G2939	1311 Bt1Gc1658	=
INTERGENIC REGION [Escherichia coli]										
HYPOTHETICAL 21.7 KD PROTEIN IN BETT-PRPR	8 100	31 38	1.00E-31	348	342	£2492902	11325-10723	Bt1G2938	1311 Bt1Gc1658	=
BBOTERI VAOS TREITARIOS ON NOBORALISTO		05 4/	0.005-00	000	4	64674478	9430-0330	ונעצטוום	1311 011001030	:
DECOMPROTER ANGROPHED AND EDUCATION OF THE PROPERTY AND EDUCATION			3.20E-28		31/	23913230 23913230	2/44-1924			. .
(ABUUU61/) YeeH [Bacillus subtilis]	_		8.40E-115	_	1087	g2415/45	8845-7751			
			3.00E-185		1743	g2337795	4882-7560			
(D8396)			9.10E-47		480	g2626822	4288-3827			=
(AJ010131) hypothetical protein [Bacillus cereus]	5 99	16 95	4.90E-16	200	577	g4584098	3447-3819	Bt1G2932	1310 Bt1Gc1650	_
(AJ010131) yfkH [Bacillus cereus]	_		7.30E-93		1398	g4584097	3248-2382		1310 Bt1Gc1650	=
[Bacillus subțilis]				•						
(Z99111) similar to heavy metal-transporting ATPase	5 100		9.10E-173	1679	1789	g2633756	269-2182	Bt1G2930	1310 Bt1Gc1650	=
CARBON STARVATION PROTEIN A [Escherichia coli]	0 99	85 60	4.70E-185	1795	2207	g2506997	13243-11163	Bt1G2929	1309 Bt1Gc1654	<u></u>
					į					:
HYPOTHETICAL 7.7 KD PROTEIN IN MRR-TSR	7 99	09 37	1.80E-09	138	131	g732103	11187-10987	Bt1G2928	1309 Bt1Gc1654	
INCDI BI MESCI PROB	t Cyrg	Ident	Prob	Score	nap Score	MCDI 81	T OSITION	Oche In	NO Contrigue	z -
NOBI di Association	%	%	BlastP-	BlastP	aat	NCRI ei	Docition	Cene Id	SEQ Cantia Id	. 2

(AE00	40 100	2.10E-69	562	624	g1787450	3965-2943	Bt1G2975	1317 Bt1Gc1661	
	•			(9				
		1.40E-57	592	575	e2497392	1-368	Bt1G2974	1317 Bt1Gc1661	
(AJ22	29 99	6.50E-21	246	205	g2687738	9651-10196	2 Bt1G2973	1316 Bt1Gc1662	
_	38 99	3.20E-12	<u>1</u> 2	146	g2634222	9654-9938	2 Bt1G2972	1316 Bt1Gc1662	
0 (X98106) zinc finger protein [Bacteriophage phig1e]	28 100	1.10E-25	291	165	g1926335	2946-3806		1316 Bt1Gc1662	
_		2.70E-15	193	183	82/0/959	08/-899			
			3	3					
(Z991	74 100	8.00E-87	868	972	g2635188	7245-7970	5 Bt1G2969	1315 Bt1Gc1656	
protein [Bacillus subtilis]					• •				
0 (Z99118) similar to sodium/proton-dependent alanine carrier	68 100	7.50E-178	1727	1757	g2635246	5649-7106	5 Bt1G2968	1315 Bt1Gc1656	
0 (Y15254) YerI protein [Bacillus subtilis]	30 100	1.70E-43	459	335	g2577967	4509-5942		1315 Bt1Gc1656	
subtilis]									
0 (D50453) spore germination protein GerKA [Bacillus	35 100	9.90E-89	886	865	g1805440	4383-2791	5 Bt1G2966	1315 Bt1Gc1656	
9 (Y09254) hypothetical protein [Bacillus cereus]	96 99	2.90E-57	589	589	g2072375	1282-917	5 Bt1G2965	1315 Bt1Gc1656	
6 (D83967) TreC [Bacillus subtilis]	56 36	6.80E-58	595	563	g2626830	1-604	5 Bt1G2964	1315 Bt1Gc1656	
pneumoniae]					٠.			•	
0 (U83667) macrolide-efflux determinant [Streptococcus	32 100	1.20E-42	451	532		11275-12479	5 Bt1G2963	1314 Bt1Gc1655	
	42 100	6.00E-105	1039	1083	g2443256	9725-11269	Bt1G2962	1314 Bt1Gc1655	
(Diamine acetyltransferase) (SAT). [Escherichia coli]							٠		
_	63 99	4.00E-60	616	642	g1742583	8769-9323	5 Bt1G2961	1314 Bt1Gc1655	
	51 99	2.40E-21	250	370	g2460264	8219-8626	Bt1G2960	1314 Bt1Gc1655	
	94 100	2.60E-246	2373	2327	g3334367	4059-5511	Bt1G2959		
protein FecE of E. coli [Bacillus subtilis]									
	46 · 100	1.20E-58	602	596	g1805452	2840-3595	5 Bt1G2958	1314 Bt1Gc1655	
permerase protein FatC of V. anguillarum [Bacillus subtilis]								•	
	33 100	2.50E-44	467	497	g1805451	1899-2843	5 Bt1G2957	1314 Bt1Gc1655	
(X8884)	39 100	8.40E-44	462	632	g1107528	821-1770	Bt1G2956	1314 Bt1Gc1655	
protein [Neisseria gonorrhoeae]					٠			•.	
(AF11	34 72	8.20E-30	330	190	g4768683	1-752	6 Bt1G2955	1314 Bt1Gc1655	
_	43 92	3.60E-226	2183	2056	g4760825	10755-6797	Bt1G2954	1313 Bt1Gc1659	
_	37 99	4.90E-17	188	271	g2635586	7542-7016	Bt1G2953	1313 Bt1Gc1659	
INTERGENIC REGION [Bacillus subtilis]									
	64 97	2.30E-112	1109	1584	g3915560	6981-5413	Bt1G2952	1313 Bt1Gc1659	
(H1) [Bacillus subtilis]								-	
TACET BLASSCIPTION	Ident Cyrg	Prob Id	Score	nap Score	. Cor Br	T OSITION	Oche Id	NO County to	
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Bt1Gc1666	Bt1Gc1666	Bt1Gc1663	Bt1Gc1663	Bt1Gc1663	-	Bt1Gc1663	Bt1Gc1663	-	Bt1Gc1663		•		1318 Bt1Gc1663		1318 BUTC1003	7.10-100	1318 Bt1Gc1663		00100100	Bt1Gc1661	Bt1Gc1661		Bt1Gc1661	Bt1Gc1661		Contig Id	
Bt1G2991	Bt1G2990	Bt1G2989	Bt1G2988	Bt1G2987		Bt1G2986	Bt1G2985		Bt1G2984				Bt1G2983		B[102982		Bt1G2981		D1102290	Bt1G2979	Bt1G2978		Bt1G2977	Bt1G2976		Gene Id	
3660-2478	2305-1	11635-12268	11003-11644	9774-11949		7487-9799	6441-7513		4843-6379				3831-4856		3293-3804	2202	1580-3292		392-1204	12204-11780	10089-11255		6523-5918	4670-5701		Position	
g3915989	g3123226	8 g3·122231		g2618864		g3122355	g1770070		g2492784				2585314		g11/0349	1170540	g1770066	٠.	G177712				g3183496	g1787450		NCBI gi	
1669	371	402	455	457		1487	927		1410			į	1235		524 ₩	3	1908		. 000	387	419		313	463	Score	nap	
·	474	457	488	544		1208	987		1441				1250		492		1880		620	406 26	380	•	³ 00	421		BlastP Score	
5.20E-154	2.10E-44	2.80E-43	1.50E-46	1.70E-52		2.10E-173	2.00E-99		1.50E-147				2.60E-127		3.6UE-4/		4.60E-194		2.90E-02	7.20E-38	4.10E-35		1.20E-26	1.90E-52		BlastP- Prob	1
	28	46	4	33		48	55		56			;	69	_	0	3	2		32	52	29		33	35		% % Ident Cvrg	
100	දු	49	100	89		100	100		100				100		99		100		100		100		100	100		Cvrg	
HYPOTHETICAL 45.3 KD PROTEIN IN PRKA-CSPB INTERGENIC REGION (ORF4) [Bacillus subtilis]	[Methanococcus jannaschii] INTERNALIN A PRECURSOR []	subtilis] HISTIDINOL DEHYDROGENASE (HDH)	ATP PHOSPHORIBOSYLTRANSFERASE [Bacillus	(AF017113) histidyl-tRNA synthetase [Bacillus subtilis]	(ISOPROPYLMALATE ISOMERASE) (ALPHA-IPM	3-ISOPROPYEMALATE DEHYDRATASE	(Z75208) 3-isopropylmalate dehydrogenase [Bacillus	SYNTHETASE) [Bacillus subtilis]	2-ISOPROPYLMALATE SYNTHASE (ALPHA-	REDUCTOISOMERASE) [Bacillus subtilis]	(ALPHA-KETO-BETA-HYDROXYLACIL	(ACETOHYDROXY-ACID ISOMEROREDUCTASE)	KETOL-ACID REDUCTOISOMERASE	SUBUNIT) (ALS) [Bacillus subtilis]	(AHAS) (ACETOHYDROXY ACID SYNTHASE SMALL	subtilis]	(Z75208) acetolactate synthase large subunit [Bacillus	[Archaeoglobus fulgidus]	AMINOTRANSFERASE (TRANSAMINASE B) (BCAT)	(AE000433) IS150 putative transposase [Escherichia coli]	(D90809) Protein AraJ precursor. [Escherichia coli]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 23.4 KD PROTEIN IN AAPA-SIGV	[Escherichia coll] (AE000218) putative dihydroxyacetone kinase (EC 2.7.1.2)		NCBI gi description	•

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	DEGV PROTEIN [Bacillus subtilis] HYPOTHETICAL 24.8 KD PROTEIN IN DEGS-TAGO	100	73	1.80E-87 1.20E-85	, ,	906 846	g729311 g3123311	8984-8145 9221-9867	Bt1G3015	1322 Bt1Gc1669 1322 Bt1Gc1669	==
	(Z99121) similar to hypothetical proteins [Bacillus subtilis]	2		2.20E-29		259	g2635881	7963-7511			=
	P54 PROTEIN PRECURSOR []	60	24	1.40E-05	122	219	g129375	7371-4824	Bt1G3013	1322 Bt1Gc1669	
	(Y14079) hypothetical protein [Bacillus subtilis]	24	47	1.00E-26	304	240	g2226145	6446-6094	Bt1G3012	1322 Bt1Gc1669	
	COMF OPERON PROTEIN 1 [Bacillus subtilis]	100	49	1.40E-114	1130	1060	g729157	6903-4620	Bt1G3011	1322 Bt1Gc1669	
	COMF OPERON PROTEIN 3 [Bacillus subtilis]	100	45	2.50E-44	467	366	g729159	4617-3916	Bt1G3010	1322 Bt1Gc1669	=
	COLD SHOCK-LIKE PROTEIN CSPC [Bacillus cereus]	98	100	5.60E-31	341	. 341	g2493763	3786-3592	Bt1G3009	1322 Bt1Gc1669	
	(ORF2) [Clostridium perfringens] HYPOTHETICAL 22.0 KD PROTEIN IN FLIT-SECA	99	66	6.50E-53	548	554	g732321	3267-2725	Bt1G3008	1322 Bt1Gc1669	
	PROTEIN HI0355 [Haemophilus influenzae Rd] HYPOTHETICAL 10.7 KD PROTEIN IN VIRR 5'REGION	99	46	1.80E-18	223	232	g3025318	9441-9154	Bt1G3007	1321 Bt1Gc1665	-
	HI0357 [Haemophilus influenzae Rd] HYPOTHETICAL ABC TRANSPORTER PERMEASE	100	33	9.20E-38	405	405	g2501390	9137-8403	Bt1G3006	1321 Bt1Gc1665	=
	PUTATIVE THIAMINE BIOSYNTHESIS PROTEIN	52	32	1.30E-22	262	173	g1175228	8874-7864	Bt1G3005	1321 Bt1Gc1665	· ==
	(U50335) ORF3 [Mycobacterium smegmatis]	100	_	8.10E-46	481	440	g1477569	7393-6641	Bt1G3004	1321 Bt1Gc1665	=
	PROTEIN CY50.01 [Mycobacterium tuberculosis] (U89796) chitinase [Bacillus thuringiensis]	100	91	4.80E-270	2597	2736	g2149596	3970-6284	Bt1G3003	1321 Bt1Gc1665	=
C)	APPC [Bacillus subtilis] HYPOTHETICAL ABC TRANSPORTER ATP-BINDING	100	43	1.20E-124	1225	1230	g1723073	9378-11166	Bt1G3001	1320 Bt1Gc1664	=
	PERMEASE PROTEIN [Helicobacter pylori J99] OLIGOPEPTIDE TRANSPORT PERMEASE PROTEIN	100	4	8.10E-46	481	623	g1168474	8440-9336	Bt1G3000	1320 Bt1Gc1664	=
	acetylmuramoyl-L-alanine amidase [Bacillus subtilis] (AE001465) DIPEPTIDE TRANSPORT SYSTEM	100	39	4.40E-57	331	574	g4154809	7540-8484	Bt1G2999	1320 Bt1Gc1664	
	PROTEIN) [Bacillus licheniformis] (Z99116) alternate gene name: yqil; similar to N-	100	39	2.80E-29	325	235	g2634853	712-1370	Bt1G2998	1320 Bt1Gc1664	
, -	S-LAYER PROTEIN PRECURSOR (SURFACE LAYER	16	49	1.00E-31	358	· 317	g1351072	1411	Bt1G2997	1320 Bt1Gc1664	
	3'REGION (ORF4) [Bacillus stearothermophilus] (Z93102) hypothetical 48.5 kd protein [Bacillus subtilis]	· 99	2	1.50E-131	1290	1401		11474-10060	Bt1G2996	1319 Bt1Gc1666	н
	HYPOTHETICAL 35.5 KD PROTEIN IN GLDA	100	42	2.20E-59	609	606	g418441	10023-9130	Bt1G2995	1319 Bt1Gc1666	-
	xylinus] (U58864) CspR [Bacillus subtilis]	99	70	1.50E-60	620	619	g1381681	9080-8610	Bt1G2994	1319 Bt1Gc1666	
•	(AF052517) c-di-GMP phosphodiesterase A [Acetobacter	58	32	3.40E-63	645	589	g3659614	9541-6407		1319 Bt1Gc1666	=
	PRKA PROTEIN [Bacillus subtilis]	100	88	3.50E-306	2938	2938	g730399	6008-4116	Bt1G2992	1319 Bt1Gc1666	=
	NCBI gi description	%Cvrg	% Ident	BlastP- Prob	BlastP Score	nap Score	NCBI gi	Position	Gene Id	ID Contig Id	NO E SEC
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[Bacillus stearothermophilus] thermostable pullulanase [Bacillus stearothermophilus] (AF008220) YtmP [Bacillus subtilis]	100	3 54 5 54	7.80E-183 9.60E-75	1774 754	1823 772	g228654 g2293206	5394-7373 7595-8403	674 Bt1G3058 674 Bt1G3059	1327 Bt1Gc1674 1327 Bt1Gc1674
[Bacillus subtilis] (AF008220) putative peptidase [Bacillus subtilis] hypothetical protein 3 - Bacillus stearothermophilus	100 67	\$ 58 7 37	7.00E-143 5.10E-37	1397 398	1400 317	g2293201 g1075814	2974-4368 4472-6397	674 Bt1G3056 674 Bt1G3057	1327 Bt1Gc1674 1327 Bt1Gc1674
INTERGENIC REGION [Bacillus subtilis] (Z99119) similar to transcriptional regulator (DeoR family)	99	5 74	1.10E-25	291	291	g2635486	2607-2389	674 Bt1G3055	1327 Bt1Gc1674
HYPOTHETICAL 17.7 KD PROTEIN IN AMYX-OPUD	99	54	2.60E-40	429	420	g3915559	1824-2291	674 Bt1G3054	1327 Bt1Gc1674
[Bacillus subtilis] (AF008220) YtgP [Bacillus subtilis]	93	5 51	2.20E-116	1147	1226	g2293198	1-1502		1327 Bt1Gc1674
influenzae Rd] (Z99120) similar to iron(III) dicitrate transport permease	100	5 71	3.00E-96	957	1030	g2635790	11918-11094 g2635790	Bt1G3052	1326 Bt1Gc1673
horikoshii] HYPOTHETICAL PROTEIN HI0912 [Haemophilus	100	5 31	1.20E-26	300	254	g1175329	10338-11069 g1175329	Bt1G3051	1326 Bt1Gc1673
synthase (8 amino-7-oxonenanoate synthase) [Pyrococcus				eş e					
(AP000001) 398aa long hypothetical 5-aminolevulinic acid	100	55	2.00E-113	1119	1057	g3256681	10121-8931	673 Bt1G3050	1326 Bt1Gc1673
(U64847) F08F3.4 gene product [Caenorhabditis elegans]	59	3 43	7.40E-68	689	592	g1458327	9997-8016	673 Bt1G3049	1326 Bt1Gc1673
(Z99114) yojO [Bacillus subtilis]	100	40	8.30E-124	1217	1155	g2634331	5450-7398	673 Bt1G3048	1326 Bt1Gc1673
(Z99114) similar to nitric-oxide reductase [Bacillus subtilis]	100	58	3.70E-89	890	· 869	g2634332	4588-5505	673 Bt1G3047	1326 Bt1Gc1673
(U88888) cardiolipin synthase [Bacillus firmus]	100	41	1.00E-109	1084	. 1063	g2952028	2967-4490	673 Bt1G3046	1326 Bt1Gc1673
(D90905) hypothetical protein [Synechocystis sp.]	65		1.50E-76	771	66.1	g1652490	3431-245	673 Bt1G3045	1326 Bt1Gc1673
(Z99117) similar to NifS protein homolog [Bacillus subtilis]	70	2	3.50E-84	843	. 841	g2635197	11391-10666	672 Bt1G3044	.1325 Bt1Gc1672
THIOURIDYLATE)-METHYLTRANSFERASE []						(
PROBABLE TRNA (5-METHYLAMINOMETHYL-2-	100		4.00E-156	1522	1509	g3122977	10538-9432		
subtilis] (Z99117) similar to hypothetical proteins [Bacillus subtilis]	100	53	2.70E-54	561	560	g2635194	9283-8666	672 Bt1G3042	1325 Bt1Gc1672
(Z99117) similar to conjugation transfer protein [Bacillus	100	2 56	3.30E-232	2240	2289	g2635193	8558-6213	672 Bt1G3041	1325 Bt1Gc1672
(Z99117) similar to hypothetical proteins [Bacillus subtilis]	100	4	3.30E-58	598	848	g2635187	5324-4266	672 Bt1G3040	1325 Bt1Gc1672
ALANYL-TRNA SYNTHETASE (ALANINETRNA	100	75	0	3311	3462	g3122886	3367-731	672 Bt1G3039	1325 Bt1Gc1672
(Z99117) similar to hypothetical proteins [Bacillus subtilis]	91		2.90E-41	438	457	g2635185	378-1	672 Bt1G3038	1325 Bt1Gc1672
[Bacillus subtilis] (Y14080) hypothetical protein [Bacillus subtilis]	40	56	5.30E-10	143	124	g2226157	11468-11326	671 Bt1G3037	1324 Bt1Gc1671
	CVF	Ident Cyrg	FTOD	Score	Score		÷		
NCBI gi description	%	%	BlastP-	BlastP	aat_	NCBI gi	Position	Id Gene Id	SEQ ID Contig Id
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(L035) (U889				130 53	g1850913 g2444121	10864-1 4731-3283			1330 1330
PERMEASE PROTEIN POTC [] (AF047044) putative transposase [Anabaena PCC7120]	25 32	5.60E-09	139 5	93	g3005554	1-307	Bt1G3082	0 Bt1Gc1677	1330
	40 100	1.90E-37	402 1	526	g1172564	16391-15627	Bt1G3081	9 Bt1Gc1675	1329
	35 100	6.90E-49	510 6	564	g1172561	15552-14512	Bt1G3080	9 Bt1Gc1675	1329
hypothetical protein 2 - Bacillus megaterium [Bacillus	31 100	1.70E-22	261 1	254	g322157	14449-13799	Bt1G3079	9 Bt1Gc1675	1329
(Z95398) unknown [Mycobacterium leprae]	28 62	7.30E-55	Ī	466	g2104606	13196-9890	Bt1G3078	9 Bt1Gc1675	1329
INTERGENIC REGION [Bacillus subtilis]									
[Bacillus subtilis] HYPOTHETICAL 33.7 KD PROTEIN IN CSPB-GLPP	46 100	3.30E-72	730 3	726	g1724014	10868-9963	Bt1G3077	9 Bt1Gc1675	1329
(Z99111) similar to hypothetical proteins from B. subtilis	51 100	6.50E-53	548 6	521	g2633768	8374-9198	Bt1G3076	9 Bt1Gc1675	1329
(Z75208) hypothetical protein [Bacillus subtilis]	68 100	8.20E-156	1519 8.2	1622	g1770025	6577-7908	Bt1G3075	9 Bt1Gc1675	1329
_	75 100	1.80E-183	1780 1.3	1847	g1770026	5171-6580		9 Bt1Gc1675	1329
_	53 50	1.00E-24	282 1	244	g1945709	334-1	Bt1G3073	9 Bt1Gc1675	1329
_		7.60E-27		282	g1787091	15716-16030	Bt1G3072	8 Bt1Gc1676	1328
coelicolor]									
[Bacillus subtilis] (AL031031) putative sensory histidine kinase [Streptomyces	35 40	1.20E-99	737 1	905	g3294236	16729-11142	Bt1G3071	8 Bt1Gc1676	1328
CHEMOTAXIS PROTEIN METHYLTRANSFERASE	26 100	7.60E-27	302 7	227	g584925	11071-10292	Bt1G3070	8 Bt1Gc1676	1328
system [Synechocystis sp.] (Z94043) hypothetical protein [Bacillus subtilis]	31 68	4.80E-41	436. 4	363	g1945716	8005-10219	Bt1G3069	8 Bt1Gc1676	1328
	37 51	2.10E-13	151 2	164	g1653076	9054-11223	Bt1G3068	8 Bt1Gc1676	1328
	53 · 100	3.00E-71		711	g3386359	7614-8411	Bt1G3067	8 Bt1Gc1676	1328
jannaschii] (AF008220) putative thioredoxin [Bacillus subtilis]	54 · 86	3.20E-28	315 3	314	g2293211	13132-13409	Bt1G3066	7 Bt1Gc1674	1327
HYPOTHETICAL PROTEIN MJ0261 [Methanococcus	27 100	3.70E-18	220 3	103	g3915930	12396-13051	Bt1G3065	7 Bt1Gc1674	1327
(AF008220) YtoP [Bacillus subtilis]	74 100	3.50E-148	1447 3.5	1446	g2293210	11185-12255	Bt1G3064	7 Bt1Gc1674	1327
_	36 99	7.40E-13	170 7	171	g2635471	11036-10719	Bt1G3063	7 Bt1Gc1674	1327
(AF008220) YmP [Bacillus subtilis]	62 100	2.20E-82	826 2	882	g2293208	9707-10474	Bt1G3062	7 Bt1Gc1674	1327
_	65 100	1.10E-66	678 1	739	g2293207	8909-9547	Bt1G3061	7 Bt1Gc1674	1327
(Z99119) ytzH [Bacillus subtilis]	54 99	1.40E-20	243 1	279	g2635475	8680-8405	Bt1G3060	7 Bt1Gc1674	1327
NCBI gi description	nt Cyrg	BlastP- % Prob Ident	BlastP Bi Score I	nap S Score	NCBI gi	Position	Gene Id	Contig Id	NO EQ

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subtilis	HYPOTHETICAL 8.8 KD PROTEIN IN SPO INTERGENIC REGION [Bacillus subtilis] TRANSCRIPTION-REPAIR COUPLING FAIRTCF) [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Ba (AF023181) low temperature requirement B pi [Listeria monocytogenes]	HYPOTHETICAL 8.8 KD PROTEIN IN SPO INTERGENIC REGION [Bacillus subtilis] TRANSCRIPTION-REPAIR COUPLING FA (TRCF) [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Ba (AF023181) low temperature requirement B p [Listeria monocytogenes] HYPOTHETICAL 56.1 KD PROTEIN IN MI INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 8.8 KD PROTEIN IN SPO INTERGENIC REGION [Bacillus subtilis] TRANSCRIPTION-REPAIR COUPLING FA (TRCF) [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Ba (AF023181) low temperature requirement B p [Listeria monocytogenes] HYPOTHETICAL 56.1 KD PROTEIN IN MF INTERGENIC REGION [Bacillus subtilis] (AB016427) thioesterase II-like protein [Bacilichenischemic]	HYPOTHETICAL 8.8 KD PROTEIN IN SPOVC-MFD INTERGENIC REGION [Bacillus subtilis] TRANSCRIPTION-REPAIR COUPLING FACTOR (TRCF) [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Bacillus subtilis] (AF023181) low temperature requirement B protein [Listeria monocytogenes] HYPOTHETICAL 56.1 KD PROTEIN IN MFD-DIVIC INTERGENIC REGION [Bacillus subtilis] (AB016427) thioesterase II-like protein [Bacillus licheniformis] GSP PROTEIN [Brevibacillus brevis] GSP PROTEIN [Brevibacillus brevis] (AF004835) tyrocidine synthetase 3 [Brevibacillus brevis] (Z99113) yoeD [Bacillus subtilis]	HYPOTHETICAL 8.8 KD PROTEIN IN SPON INTERGENIC REGION [Bacillus subtilis] TRANSCRIPTION-REPAIR COUPLING FAC (TRCF) [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Bac (AF023181) low temperature requirement B proflisteria monocytogenes] HYPOTHETICAL 56.1 KD PROTEIN IN MF1 INTERGENIC REGION [Bacillus subtilis] (AB016427) thioesterase II-like protein [Bacillicheniformis] GSP PROTEIN [Brevibacillus brevis] (AF004835) tyrocidine synthetase 3 [Brevibac (Z99113) yoeD [Bacillus subtilis] SPORE COAT PROTEIN B [Bacillus subtilis]	HYPOTHETICAL 8.8 KD PROTEIN IN SPO INTERGENIC REGION [Bacillus subtilis] TRANSCRIPTION-REPAIR COUPLING FA (TRCF) [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Ba (AF023181) low temperature requirement B p [Listeria monocytogenes] HYPOTHETICAL 56.1 KD PROTEIN IN MF INTERGENIC REGION [Bacillus subtilis] (AB016427) thioesterase II-like protein [Bacilicheniformis] GSP PROTEIN [Brevibacillus brevis] (AF004835) tyrocidine synthetase 3 [Brevibacillus subtilis] SPORE COAT PROTEIN B [Bacillus subtilis] SPORE COAT PROTEIN B [Bacillus subtilis]	HYPOTHETICAL 8.8 KD PROTEIN IN SPC INTERGENIC REGION [Bacillus subtilis] TRANSCRIPTION-REPAIR COUPLING FA (TRCF) [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Bacillus amonocytogenes] HYPOTHETICAL 56.1 KD PROTEIN IN MINTERGENIC REGION [Bacillus subtilis] (AB016427) thioesterase II-like protein [Bacilicheniformis] GSP PROTEIN [Brevibacillus brevis] (AF004835) tyrocidine synthetase 3 [Breviba (Z99113) yoeD [Bacillus subtilis] SPORE COAT PROTEIN B [Bacillus subtilis] (Z99111) similar to two-component sensor his [Bacillus subtilis]	HYPOTHETICAL 8.8 KD PROTEIN IN SPOVC-MFD INTERGENIC REGION [Bacillus subtilis] TRANSCRIPTION-REPAIR COUPLING FACTOR (TRCF) [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Bacillus subtilis] (AF023181) low temperature requirement B protein [Listeria monocytogenes] HYPOTHETICAL 56.1 KD PROTEIN IN MFD-DIVIC INTERGENIC REGION [Bacillus subtilis] (AB016427) thioesterase II-like protein [Bacillus licheniformis] GSP PROTEIN [Brevibacillus brevis] (AF004835) tyrocidine synthetase 3 [Brevibacillus brevis] (AF004835) tyrocidine synthetase 3 [Brevibacillus brevis] (Z99113) yoeD [Bacillus subtilis] (Z99767) ywrJ [Bacillus subtilis] (Z99767) ywrJ [Bacillus subtilis] (Z99111) similar to two-component sensor histidine kinase [Bacillus subtilis] (AL022003) hypothetical protein Rv1716 [Mycobacterium tuberculosis]	HYPOTHETICAL 8.8 KD PROTEIN IN SPOINTERGENIC REGION [Bacillus subtilis] TRANSCRIPTION-REPAIR COUPLING FA (TRCF) [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Bacillus subtilis] KAF023181) low temperature requirement B p [Listeria monocytogenes] HYPOTHETICAL 56.1 KD PROTEIN IN MINTERGENIC REGION [Bacillus subtilis] (AB016427) thioesterase II-like protein [Bacilicheniformis] (AB016427) thioesterase II-like protein [Bacilicheniformis] GSP PROTEIN [Brevibacillus brevis] (AF004835) tyrocidine synthetase 3 [Breviba (Z99113) yoeD [Bacillus subtilis] SPORE COAT PROTEIN B [Bacillus subtilis] (Z93767) ywrJ [Bacillus subtilis] (Z99111) similar to two-component sensor hin [Bacillus subtilis] (AL022003) hypothetical protein Rv1716 [Muberculosis] (D50453) ycgL [Bacillus subtilis]	HYPOTHETICAL 8.8 KD PROTEIN IN SPONTERGENIC REGION [Bacillus subtilis] TRANSCRIPTION-REPAIR COUPLING FA (TRCF) [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [B (AF023181) low temperature requirement B 1 [Listeria monocytogenes] HYPOTHETICAL 56.1 KD PROTEIN IN M INTERGENIC REGION [Bacillus subtilis] (AB016427) thioesterase II-like protein [Bacilcheniformis] (AF004835) tyrocidine synthetase 3 [Brevibacilcheniformis] (AF004835) yoeD [Bacillus subtilis] (Z99111) similar to two-component sensor hi [Bacillus subtilis] (Z99111) similar to two-component sensor hi [Bacillus subtilis] (AL022003) hypothetical protein Rv1716 [Iv tuberculosis] (AF009224) benzoate transport protein [Acid ADP1]	HYPOTHETICAL 8.8 KD PROTEIN IN SPOVC-MFD INTERGENIC REGION [Bacillus subtilis] TRANSCRIPTION-REPAIR COUPLING FACTOR (TRCF) [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Bacillus subtilis] (AF023181) low temperature requirement B protein [Listeria monocytogenes] HYPOTHETICAL 56.1 KD PROTEIN IN MFD-DIVIC INTERGENIC REGION [Bacillus subtilis] (AB016427) thioesterase II-like protein [Bacillus licheniformis] GSP PROTEIN [Brevibacillus brevis] (AF004835) tyrocidine synthetase 3 [Brevibacillus brevis] (AF004835) yoeD [Bacillus subtilis] (Z99111) similar to two-component sensor histidine kinase [Bacillus subtilis] (Z99111) similar to two-component sensor histidine kinase [Bacillus subtilis] (AL022003) hypothetical protein Rv1716 [Mycobacteriun tuberculosis] (AF009224) benzoate transport protein [Acinetobacter sp. ADP1] (Y14084) hypothetical protein [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis] TRANSCRIPTION-REPAIR COUPLING FACT (TRCF) [Bacillus subtilis] STAGE V SPORULATION PROTEIN T [Bacill (AF023181) low temperature requirement B prote [Listeria monocytogenes] HYPOTHETICAL 56.1 KD PROTEIN IN MFD-INTERGENIC REGION [Bacillus subtilis] (AB016427) thioesterase II-like protein [Bacillus licheniformis] (AB016427) thioesterase II-like protein [Bacillus licheniformis] (AF004835) tyrocidine synthetase 3 [Brevibacille (AF004835) tyrocidine synthetase 3 [Brevibacille (Z99113) yoeD [Bacillus subtilis] (Z99111) similar to two-component sensor histidi [Bacillus subtilis] (Z99111) similar to two-component sensor histidi [Bacillus subtilis] (AL022003) hypothetical protein Rv1716 [Mycc tuberculosis] (AF009224) benzoate transport protein [Acineto ADP1] (X14084) hypothetical protein [Bacillus subtilis] ORF 1S231C [Bacillus thuringiensis]
	Bt1Gc1667 Bt1G3091 7069-7602 g586883 727 727 7.00E-72 77 99 STAGE Bt1Gc1667 Bt1G3092 7851-9440 g4090864 745 624 5.70E-61 36 100 (AF023	Bt1Gc1667 Bt1G3091 7069-7602 g586883 727 727 7.00E-72 77 99 STAGE Bt1Gc1667 Bt1G3092 7851-9440 g4090864 745 624 5.70E-61 36 100 (AF023 Bt1Gc1667 Bt1G3093 9450-10287 g586885 912 917 5.10E-92 61 57 HYPOTINTER	Bt1Gc1667 Bt1G3091 7069-7602 g586883 727 727 7.00E-72 77 99 STAGE Bt1Gc1667 Bt1G3092 7851-9440 g4090864 745 624 5.70E-61 36 100 (AF023 Bt1Gc1667 Bt1G3093 9450-10287 g586885 912 917 5.10E-92 61 57 HYPOTINTER Bt1Gc1668 Bt1G3094 1270-568 g4126668 491 512 4.20E-49 42 100 (AB016	Bt1Gc1667 Bt1G3091 7069-7602 g586883 727 727 7.00E-72 77 99 STAGE Bt1Gc1667 Bt1G3092 7851-9440 g4090864 745 624 5.70E-61 36 100 (AF023 Bt1Gc1667 Bt1G3093 9450-10287 g586885 912 917 5.10E-92 61 57 HYPO3 Bt1Gc1668 Bt1G3094 1270-568 g4126668 491 512 4.20E-49 42 100 (AB010 Bt1Gc1668 Bt1G3095 1988-1275 g1170075 434 497 1.60E-47 42 100 GSP PR Bt1Gc1668 Bt1G3096 17853-1 g2623773 6940 7957 0 39 67 (AF004 Bt1Gc1679 Bt1G3097 1-215 g2634223 257 259 2.70E-22 68 93 (29911	Bt1Gc1667 Bt1G3091 7069-7602 g586883 727 727 7,00E-72 77 99 STAGE Bt1Gc1667 Bt1G3092 7851-9440 g4090864 745 624 5.70E-61 36 100 (AF023 Bt1Gc1667 Bt1G3093 9450-10287 g586885 912 917 5.10E-92 61 57 HYPO3 Bt1Gc1668 Bt1G3094 1270-568 g4126668 491 512 4.20E-49 42 100 (AB016 Bt1Gc1668 Bt1G3095 1988-1275 g1170075 434 497 1.60E-47 42 100 GSP PF Bt1Gc1668 Bt1G3096 17853-1 g2623773 6940 7957 0 39 67 (AF004 Bt1Gc1679 Bt1G3098 55-3110 g2815495 261 371 3.70E-34 35 67 SPORE	Bt1Gc1667 Bt1G3091 7069-7602 g586883 727 727 7,00E-72 77 99 STAGE Bt1Gc1667 Bt1G3092 7851-9440 g4090864 745 624 5,70E-61 36 100 (AF023 Bt1Gc1667 Bt1G3093 9450-10287 g586885 912 917 5,10E-92 61 57 HYPO3 Bt1Gc1668 Bt1G3094 1270-568 g4126668 491 512 4,20E-49 42 100 (AB016 Bt1Gc1668 Bt1G3095 1988-1275 g1170075 434 497 1,60E-47 42 100 GSP PF Bt1Gc1668 Bt1G3096 17853-1 g2623773 6940 7957 0 39 67 (AF004 Bt1Gc1679 Bt1G3098 55-3110 g2815495 261 371 3,70E-34 35 67 SPORE Bt1Gc1679 Bt1G3099 1595-2260 g1929337 468 477 2,20E-45 40 100 (29376 <td>Bt1Gc1667 Bt1G3091 7069-7602 g586883 727 727 7.00E-72 77 99 STAGE Bt1Gc1667 Bt1G3092 7851-9440 g4090864 745 624 5.70E-61 36 100 (AF023 Bt1Gc1667 Bt1G3093 9450-10287 g586885 912 917 5.10E-92 61 57 HYPO3 Bt1Gc1668 Bt1G3094 1270-568 g4126668 491 512 4.20E-49 42 100 (AB016 Bt1Gc1668 Bt1G3095 1988-1275 g1170075 434 497 1.60E-47 42 100 GSP PR Bt1Gc1668 Bt1G3096 17853-1 g2623773 6940 7957 0 39 67 (AF004 Bt1Gc1679 Bt1G3098 55-3110 g2815495 261 371 3.70E-34 35 67 SPORE Bt1Gc1679 Bt1G3100 4439-5156 g2633724 420 487 4.30E-45 40 100 (Z9376 <td>Bt1Gc1667 Bt1G3091 7069-7602 g586883 727 727 7.00E-72 77 99 STAGE Bt1Gc1667 Bt1G3092 7851-9440 g4090864 745 624 5.70E-61 36 100 (AF023 Bt1Gc1667 Bt1G3093 9450-10287 g586885 912 917 5.10E-92 61 57 HYPO3 Bt1Gc1668 Bt1G3094 1270-568 g4126668 491 512 4.20E-49 42 100 (AB01t Bt1Gc1668 Bt1G3095 1988-1275 g1170075 434 497 1.60E-47 42 100 GSP PH Bt1Gc1668 Bt1G3096 17853-1 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SEQ ID Contig Id Gene Id	Position	NCBI gi	nap	BlastP Score	BlastP- Prob	% Ident	Cyrg	NCBI gi description
1334 Bt1Gc1682 Bt1G3107	3742-833	g2226128	932	1121	1.20E-113	29	100	(Y14078) Hypothetical protein [Bacillus subtilis]
Bt1Gc1682	3742-833	g2226128	932	1121	1.20E-113	29	100	(Y14078) Hypothetical protein [Bacillus subtilis]
	4974-3742	g2226129	801	819	1.20E-81		100	(Y14078) Hypothetical protein [Bacillus subtilis
Bt1Gc1682	5909-6043	g2443232	151:	151	7.60E-11	62	98	(D86417) YflJ [Bacillus subtilis]
• •	7157-6174	g1731309	817	814	4.20E-81	47	100	HYPOTHETICAL 37.2 KD PROTEIN IN IDH-DEOR
		>))	·			•	INTERGENIC REGION [Bacillus subtilis]
Bt1Gc1682	7366-10511	g3242253	200	335	1.30E-28		14	(Z83857) ppsC [Mycobacterium tuberculosis]
1334 Btl'Gc1682 BtlG3113	8427-10077	g2961083	255	329	1.00E-29	37	85	(AF033674) unknown [Pseudomonas marginalis pv.
1334 Bt1Gc1682 Bt1G3114	8433-9485	g731090	159	237	5.80E-20	25	100	UXU OPERON TRANSCRIPTIONAL REGULATOR
D+1C-1692	0106 10240	~2506000	300	3	6 000		3	[Escherichia coli]
1334 BUIGE1082 BUIGS113	9190-10349	8200029	380	309	6.UUE-34	2/	Ī	CYANA IE I KANSPORT PROTEIN CYNX [symmetic construct]
1335 Bt1Gc1680 Bt1G3116	1-257	g225559	409	425	7.00E-40	99	18	ORF IS231C [Bacillus thuringiensis]
1335 Bt1Gc1680 Bt1G3117	950-3340	g3328629	140	228	5.30E-19	30	78	(AE001295) hypothetical protein [Chlamydia trachomatis]
1335 Bt1Gc1680 Bt1G3118	2938-2183	g1706579	134	155	1.70E-10		100	PROTEIN ECSC [Bacillus subtilis]
1335 Bt1Gc1680 Bt1G3119	5504-4320	g584719	1687	1604	8.10E-165		100	ACETATE KINASE (ACETOKINASE) [Bacillus subtilis]
1335 Bt1Gc1680 Bt1G3120	6793-5807	g3916031	856	879	5.40E-88	52	100	HYPOTHETICAL 37.4 KD PROTEIN IN ACKA-SSPA
								INTERGENIC REGION [Bacillus subtilis]
1335 Bt1Gc1680 Bt1G3121	7489-6989	g2293238	590	573	1.50E-55	68	99	(AF008220) YtgI [Bacillus subtilis]
1335 Bt1Gc1680 Bt1G3122	8022-7600	g2293237	370	284	6.10E-25	50	99	(AF008220) YtfJ [Bacillus subtilis]
1335 Bt1Gc1680 Bt1G3123	8876-9427	g2462088	857	821	7.60E-82	91	99	(Y11170) BC541A protein [Bacillus cereus]
1335 Bt1Gc1680 Bt1G3124	11482-9920	g2293233	1275	1262	1.40E-128	48.	100	(AF008220) YtcJ [Bacillus subtilis]
Bt1Gc1678	939-292	g2337808	858	849	8.20E-85		100	(Y13937) putative CfxE protein [Bacillus subtilis]
Bt1Gc1678	1823-945	g2337807	995	1016	1.70E-102	2	100	(Y13937) YloQ protein [Bacillus subtilis]
1336 Bt1Gc1678 Bt1G3·127	4043-2070	g2337806	1553	1550	4.30E-159	51	100	(Y13937) putative Pkn2 protein [Bacillus subtilis]
Bt1Gc1678	4804-4046	g2337805	647	629	1.70E-61	49	100	(Y13937) putative Ptc1 protein [Bacillus subtilis]
1336 Bt1Gc1678 Bt1G3129	5900-4812	g3287951	1537	1537	1.00E-157	78	100	HYPOTHETICAL 41.6 KD PROTEIN IN FMT-SPOVM
			!					INTERGENIC REGION [Bacillus subtilis]
1336 Bt1Gc1678 Bt1G3130	7239-5905	g3915867	1385	1400	3.40E-143	59	100	SUN PROTEIN [Bacillus subtilis]
.0336 Bt1Gc1678 Bt1G3131	8177-7227	g2337802	1145	1152	6.40E-117	69	100	(Y13937) putative Fmt protein [Bacillus subtilis]
1336 Bt1Gc1678 Bt1G3132	8674-8192	g2499922	451	484	3.90E-46	57	. 99	POLYPEPTIDE DEFORMYLASE (PDF)
								(FORMYLMETHIONINE DEFORMYLASE) [Bacillus subrilis]
								subtilis

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	1338 Bt1Gc1684 Bt1G3150	Bt1Gc1684	Bt1Gc1684		1338 Bt1Gc1684 Bt1G3146		1338 Bt1Gc1684			1338 Bt1Gc1684				Bt1Gc1684	Bt1Gc1684		Bt1Gc1684	Bt1Gc1681		1337 Bt1Gc1681		Bt1Gc1681	1336 Bt1Gc1678	Bt1Gc1678	Bt1Gc1678	Bt1Gc1678	BUGCIO/8		Contig Id
	Bt1G3150	Bt1G3149	Bt1G3147		Bt1G3146		Bt1G3145			Bt1G3144				Bt1G3143	Bt1G3142		Bt1G3141	Bt1G3140		Bt1G3139		Bt1G3138	Bt1G3137	Bt1G3136	Bt1G3135	Bt1G3134	BUUSISS		Gene Id
	8980-11622	7446-8478	6378-7022		4986-6275		4000-4971			3215-3988				2289-3230	1452-2273		91-2455	8755-7827		4411-3218		2360-3145	16875-14669	14518-13646	13418-12684	12293-11085	11091-8689		Position
	g68538	g586918	g421533	`	g399784		g399870		· (e399875			0	g122839	93172392	(g122819	g1817536		g1176955		g2497492			-	g2337799	g3183349		NCBI gi
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	0	8.30E-53	5.50E-30		6.10E-183		1.40E-130			2.90E-18		•		3.70E-105	6.30F-78		1.90E-156	5.20E-67		1.10E-68		9.80E-50	1.70E-249	2.00E-58	1.80E-82	1.20E-133	8.60E-248	2	BlastP- Prob
	78	30			80		75	٠.		<u>ω</u>			. (65	26	•	66	45		37		40	70	45	68	63	9	;	% Ident
	100	ž.,	37 37		100	•	100			100				_	<u>=</u>			100	•	100		100	8 3	100	100	100	100		% Cvrg
	3'REGION (ORF2) [Bacillus subtilis] valinetRNA ligase (EC 6.1.1.9) - Bacillus stearothermophilus []	HYPOTHETICAL 40.6 KD PROTEIN IN SPOVID		(GSA) (GLUTAMATE-I-SEMIALDEHYDE AMINOTRANSFERASE) (GSA-AT) [Bacillus subtilis]	GLUTAMATE-1-SEMIALDEHYDE 2,1-AMINOMUTASE	(PORPHOBILINOGEN SYNTHASE) (ALAD) (ALADH) [Bacillus subtilis]	(CYCLIZING)) [Bacillus subtilis] DELTA-AMINOLEVULINIC ACID DEHYDRATASE	(HYDROXYMETHYLBILANE HYDROLYASE	(UROPORPHYRINOGEN-III COSYNTHETASE)	UROPORPHYRINOGEN-III SYNTHASE (UROS)	subtilis]	(PRE-UROPORPHYRINOGEN SYNTHASE) (Bacillus	(HYDROXYMETHYLBILANE SYNTHASE) (HMBS)	PORPHOBILINOGEN DEAMINASE (PBG)	(AF064058) Hem X [Bacillus stearothermonhilus]	subtilis]	GLUTAMYL-TRNA REDUCTASE (GLUTR) [Bacillus	(D78508) Yfil [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis].	[Synechocyshs sp.] HYPOTHETICAL 45.8 KD PROTEIN IN ACDA-NARI	MONOPHOSPHATE KINASE) (UMP KINASE)	URIDYLATE KINASE (UK) (URIDINE	(Y13937) putative PacL protein [Bacillus subtilis]	(Y13937) YloC protein [Bacillus subtilis]	(Y13937) putative Gmk protein [Bacillus subtilis]	(Y13937) putative Dfp protein [Bacillus subtilis]	PRIMOSOMAL PROTEIN N' (REPLICATION FACTOR)		NCBI gi description

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(U86377) (p)ppGpp synthetase [Bacillus subtilis]	100	77	5.00E-307	2946	2972	g2739435	3305-1125	Bt1G3172	1342 Bt1Gc1689	_
(Z99118) similar to hypothetical proteins [Bacillus subtilis]	99	2	6.30E-39	416	415	g2635223	1024-668		1342 Bt1Gc1689	_
(U67998) unknown [Sinorhizobium meliloti]	100	33	6.00E-82	822	711	g1658371	12969-11188	6 Bt1G3170	1341 Bt1Gc1686	_
(Y09476) YisY [Bacillus subtilis]	100	.41	1.60E-58	601	584	g2145391	10189-11001	6 Bt1G3169	1341 Bt1Gc1686	_
[Bacillus subtilis]										
(Z99111) similar to two-component sensor histidine kinase	31	40	1.20E-39	429	354	g2633724	14867-7457	Bt1G3168	1341 Bt1Gc1686	
(AF017113) HPr(Ser-P) phosphatase [Bacillus subtilis]	100	33	2.00E-26	298	291	g2618860	6116-6766	Bt1G3167	1341 Bt1Gc1686	_
regulator [Bacillus subtilis]				•		•				
(Z83337) highly similar to phosphotransferase system	100	41	8.80E-49	509	502	g1763710	6019-5246	Bt1G3166	1341 Bt1Gc1686	_
(AF017113) HPr(Ser-P) phosphatase [Bacillus subtilis]	100		1.10E-18	225	169	g2618860	5258-4623	Bt1G3165	1341 Bt1Gc1686	_
INTERGENIC REGION [Bacillus subtilis]										
HYPOTHETICAL 11.8 KD PROTEIN IN GLNQ-ANSR	99	31	2.50E-12	165	148	g1731088	3300-2983	Bt1G3164	1341 Bt1Gc1686	_
PRODUCT IN E. COLI. [Bacillus subtilis]			٠							
(AB001488) FUNCTION UNKNOWN, SIMILAR	100	24	1.20E-14	191	209	g1881343	2679-1759	Bt1G3163	1341 Bt1Gc1686	_
[Bacillus subtilis]										
PHOSPHOENOLPYRUVATE CARBOXYKINASE	32	67	1.70E-61	629	611	g3123253	1-519	Bt1G3162	1341 Bt1Gc1686	_
ELEMENT IS232 [Insertion sequence IS232]		٠								
TRANSPOSASE FOR INSERTION SEQUENCE	100	100	2.90E-231	2231	2231	g2497382	2590-1298	Bt1G3161	1340 Bt1Gc1688	=
TN4556 [Transposon Tn4556]										
HYPOTHETICAL 37.1 KD PROTEIN IN TRANSPOSON	29	32	1.40E-08	136	. 74	g141450	165-2224	Bt1G3160	1340 Bt1Gc1688	_
influenzae Rdl						(
HYPOTHETICAL PROTEIN HI1455 [Haemophilus	78	4	1.50E-62	639	554	g1351814	13116-12280	Bt1G3159	1339 Bt1Gc1683	_
subtilis]				•		•				
(Z99107) similar to acriflavin resistance protein [Bacillus	100	ස	3.20E-256	2412	3348	g2632985	9088-12252	Bt1G3158	1339 Bt1Gc1683	_
[Bacillus subtilis]										
(Z99120) similar to hypothetical proteins from B. subtilis	100	38	1.00E-45	480	490	g2635800	8986-8114	Bt1G3157	1339 Bt1Gc1683	<u> </u>
(D86417) YfmM [Bacillus subtilis]	100	56	3.60E-146	1428	1535	g2443256	6506-8056	Bt1G3156	1339 Bt1Gc1683	_
CSBB PROTEIN [Bacillus subtilis]	100	55	3.70E-89	890	970	g2498262	5402-6829	Bt1G3155	1339 Bt1Gc1683	_
(Z99108) yfhO [Bacillus subtilis]	100	28	5.50E-49	529	610	g2633184	2792-5287	Bt1G3154	1339 Bt1Gc1683	_
(Z99108) yfhO [Bacillus subtilis]	100	28	5.50E-49	529	610	g2633184	2792-5287	Bt1G3153	1339 Bt1Gc1683	_
[Escherichia coli]		٠				-				
(AF019746) putative beta1,3-glucosyltransferase WaaV	. 85	23	1.50E-23	271	145	g3821837	1-840	Bt1G3152	1339 Bt1Gc1683	<u></u>
[Bacillus subtilis]				-		\$ e				
FOLYLPOLYGLUTAMATE SYNTHASE (FOLYLPOLY-	100	49	1.70E-107	1063	1057	g585155	11719-13008	Bt1G3151	1338 Bt1Gc1684	. 📇
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	Bt1G3187 6009-6971	Bt1G3186 3971-5806		Bt1G3185 2655-4043	Bt1G3184 1531-2589	سِر		Bt1G3182 15053-14131	Bt1G3181 12623-13'							33180 10573-120		Bt1G3179 1915-2685			Bt1G3177 10570-12123	Bt1G3176 9825-7606		D(1031/3 /40/-0320		Bt1G3174 6415-4067	Bt1G3173 4028-3519		Come we
, ;	71 g1176954	06 g1881268		43 g3024202	89 g2829831			131 g3183453	12623-13728 g479960				-			003 g2494077		85 g1177019		-		06 g3220156		20 B1001334		67 g2635226	19 g2580524		9
	548 548	1765 1616		1276 1321	1281 1225			409 457	369 497	··.	· ·	•				1323 1317		893 893	•		1492 1348	2085 1942	`	J/1 490		2077 - 2013	718 663	Score	
	6.50E-53	4.30E-166		7.90E-135	1.20E-124	6.90E-65		2.80E-43	1.60E-47				•			2.10E-134		1.80E-89		1.00E-45	1.10E-137	1.20E-200		7.10E-4/	0 101.47	3.70E-208	4.20E-65	Fron 10	
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5'REGION [Bacillus subtilis]	HOMOLOG. [Bacillus subtilis] 100 HYPOTHETICAL 36.9 KD PROTEIN IN ACDA	ENZYME) [Bacillus subtilis] 100 (AB001488) ATP-DEPENDENT RNA HELICASE DEAD	ALANYL LIGASE (UDP-MURNAC-PENTAPEPTIDE SYNTHETASE) (D-ALANYL-D-ALANINE-ADDING	ALANYLALANINE SYNTHETASE) [Bacillus Submis] 100 UDP-N-ACETYLMURAMOYLALANYL-D-	100 D-ALANINE-D-ALANINE LIGASE (D-		INTERGENIC REGION [Bacillus subtilis]	43 HYPOTHETICAL 38.9 KD PROTEIN IN DAT-SPO0E	56 NADH oxidase - Thermoanaerobacter brockii []	DEHYDROGENASE) [Streptococcus mutans]	DEHYDROGENASE (NADP+)) (TRIOSEPHOSPHATE	(GLYCERALDEHYDE-3-PHOSPHATE	PHOSPHATE DEHYDROGENASE)	PHOSPHORYLATING GLYCERALDEHYDE 3-	PHOSPHATE DEHYDROGENASE (NON-	100 NADP-DEPENDENT GLYCERALDEHYDE-3-	INTERGENIC REGION [Bacillus subtilis]	100 HYPOTHETICAL OXIDOREDUCTASE IN PEPT-KATB	[Bacill	(Z991		100 (AF024506) SecDF protein [Bacillus subtilis]	[Bacillus subtilis]	PRODITOT IN SYNECHOOVETIC AND F COLL		100 (Z99118) similar to single-strand DNA-specific exonuclease	99 (U86377) adenine phosphoribosyltransferase; Apt [Bacillus	a.	

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·.	Bt1G3207	Bt1G3206) Bt1G3205	Bt1G3204) Bt1G3203		_	Bt1G3200	Bt1G3199	Bt1G3198	Bt1G3197) Bt1G3196		Bt1G3195	Bt1G3194	٠.			Bt1G3192		Bt1G3190		Bt1G3189			Bt1G3188		Gene Id	
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	1133-663 4374-5660	377-1	14417-13724	724-13410	13444-12646	12653-11844	8745-9218	6515-6985	6386-6162	5590-6150	5362-3982	4017-3001		697-1	11411-12719			10984-11331	10692-10976	9220-10392	7971-9093		7630-7986			8412-6973		Position	
	g2983147 g2829488	g3913706	g2293298	13724-13410 g2293297	g2492576			g2293157	g2293294	g2293156	g2829796	g3928722		g1881335	g2632778		. (g2506170	g1881273		g1881272	•	•	g1881271		NCBI gi	
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	4.40E-38 2.80E-178	5.00E-46	3.10E-62	3.30E-26	3.40E-47	1.50E-62	1.20E-49	3.10E-69	1.00E-29	1.40E-59	. 2.90E-98	1.70E-45		1.30E-70	6.20E-135			4.60E-52	5.40E-17	9.40E-116	7.30E-93		4.80E-25			1.60E-17	rrop	BlastP-	Table 1
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2 (GSA) (GLUTAMATE-1-SEMIALDEHYDE AMINOTRANSFERASE) (GSA-AT) [Bacillus subtilis]	3 [Bacillus subtilis] (AE000692) hypothetical protein [Aquifex aeolicus] GLUTAMATE-1-SEMIALDEHYDE 2,1-AMINOMUTASE	subtilis] FERRIC UPTAKE REGULATION PROTEIN HOMOLOG	(AF008220) putative transcription regulator [Bacillus	(AF008220) YtlB· [Bacillus subtilis]	HYPOTHETICAL ABC TRANSPORTER ATP-BINDING	(AF008220) YtlD [Bacillus subtilis]	(AF008220) YtkD [Bacillus subtilis]	(AF008220) YtjB [Bacillus subtilis]	(AF008220) YtjA [Bacillus subtilis]	[Bacillus subtilis] (AF008220) YtiB [Bacillus subtilis]	[Streptomyces coelicolor] CYTOCHROME D UBIQUINOL OXIDASE SUBUNIT I	(AL034355) putative cytochrome oxidase subunit II	SUBTILIS. [Bacillus subtilis]	(AB001488) SIMILAR TO YQFU, YXKD, YITB OF B.	(Z99106) similar to hypothetical proteins [Bacillus subtilis]	AND STAPHYLOCOCCUS AUREUS. [Bacillus subtilis]	PRODUCT IN MYCOBACTERIUM TUBERCUI OSIS	(AB001488) FUNCTION UNKNOWN, SIMILAR	(AB001488) FUNCTION UNKNOWN. [Bacillus subtilis]	ALANINE RACEMASE []	(AB001488) FUNCTION UNKNOWN. [Bacillus subtilis]	PRODUCT IN E. COLI AND MYCOPLASMA	(AB001488) FUNCTION UNKNOWN, SIMILAR	SYNECHOCYSTIS. [Bacillus subtilis]	PRODUCT IN H. INFLUENZAE AND	(AB001488) FUNCTION UNKNOWN, SIMILAR		NCBI gi description	

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subtilis] dihydropteroate pyrophosphorylase - Bacillus subtilis	100	62	9.40E-84	839	903	g2126927	5039-4186	Bt1G3228	1348 Bt1Gc1694 Bt1G3228	1348
PYROPHOSPHOKINASE) (PPPK) [Bacillus subtilis] DIHYDRONEOPTERIN ALDOLASE (DHNA) [Bacillus	99	62	8.30E-37	396	439	g141435	4190-3831	Bt1G3227	BtlGc1694	1348
HYDROXYMETHYLPTERIN-PYROPHOSPHOKINASE)						·				
PYROPHOSPHOKINASE (7,8-DIHYDRO-6-								•		
2-AMINO-4-HYDROXY-6-	99	59	3.80E-48	. 503	516	g585263	3831-3319	Bt1G3226	Bt1Gc1694	1348
INTERGENIC REGION [Bacillus subtilis]					:					5
HYPOTHETICAL 37.1 KD PROTEIN IN FOLK-LYSS	100	81	1.30E-143	1404	1395	g586896	3137-2142	Bt1G3225	Bt1Gc1694	1348
LIGASE) (LYSRS) [Bacillus subtilis]	•									
LYSYL-TRNA SYNTHETASE (LYSINETRNA	100	80	3.20E-209	2023	2144	g586058	1985-489	Bt1G3224	Bt1Gc1694	1348
(Z98682) YlbB protein [Bacillus subtilis]	99	49	6.60E-28	312	350	g2339998	15684-15241	Bt1G3223	Bt1Gc1693	1347
(AE000754) lipoprotein NlpD fragment [Aquifex aeolicus]	52	4	6.70E-19	227	142	g2984049	12863-13192	Bt1G3222	Bt1Gc1693	1347
milari 26606]								1		;
PROTEIN MJ0412 [Methanococcus jannaschii] (AE000633) transcriptional regulator (ren A) [Helicobacter		47	1.50F-44	469	562	p2314455	11422-10772	Bt1G3221	Bt1Gc1693	1347
pneumoniae] HYPOTHETICAL ABC TRANSPORTER ATP-BINDING	100	39	4.20E-42	446	391	g2492576	11440-10027	Bt1G3220	Bt1Gc1693	1347
(AE001681) ABC transporter permease [Chlamydia	100	26	3.10E-46	485	434	g4377342	10003-7522	Bt1G3219	Bt1Gc1693	1347
REGULATORY PROTEIN TENI [Bacillus subtilis]	100	43	3.60E-43	456	446	g135587	8242-7628	Bt1G3218	Bt1Gc1693	1347
(Z99110) similar to sarcosine oxidase [Bacillus subtilis]	100	33	6.00E-50	520	524	g2633521	7635-6510	Bt1G3217	Bt1Gc1693	1347
(Z99110) similar to thiamin biosynthesis [Bacillus subtilis]	100	80	1.10E-103	1027	1027	g2633523	6310-5543	Bt1G3216	Bt1Gc1693	1347
[Bacillus subtilis] (Z99110) similar to thiamin biosynthesis [Bacillus subtilis]	100	58	2.30E-103	1024	985	g2633524	5541-4525	Bt1G3215	Bt1Gc1693	1347
PHOSPHOHYDROLASE [Clostridium acetobutylicum] (Z99110) similar to phosphomethylpyrimidine kinase	100	56	3.20E-60	617	788	g2633525	4506-3691	Bt1G3214	Bt1Gc1693	1347
large subunit [Corynebacterium glutamicum] POTASSIUM-TRANSPORTING ATPASE A CHAIN (ATP	100	53	6.90E-136	1331	1433	g3121769	2027-366	Bt1G3213	Bt1Gc1693	1347
aeolicus] (AB024708) glutamine 2-oxoglutarate aminotransferase	100	29	1.80E-125	657	812	g4521156	5322-10311	Bt1G3212	Bt1Gc1691	1346
large subunit [Corynebacterium glutamicum] (AE000746) glutamate synthase large subunit [Aquifex	100	29	7.10E-129	908	874	g2983926	5916-10331	Bt1G3211	Bt1Gc1691	1346
(AB024708) glutamine 2-oxoglutarate aminotransferase	100	29	1.80E-125	657	812	g4521156	5322-10311	Bt1G3210	Bt1Gc1691	1346
NCBI gi description	% Cvrg	% Ident	BlastP- Prob	BlastP Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	SEQ NO

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SEQ ID Contig Id	1348 Bt1Gc1694	1348 Bt1Gc1694		1348 Bt1Gc1694	1348 Bt1Gc1694		1348 Bt1Gc1694	1348 Bt1Gc1694		1348 811001054	1349 Bt1Gc1697	1349 Bt1Gc1697	1349 Bt1Gc1697		1349 Bt1Gc1697	1349 Bt1Gc1697	1349 Bt1Gc1697	1349 Bt1Gc1697	1350 Bt1Gc1696	1350 Bt1Gc1696	
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Gene Id	Bt1G3229	Bt1G3230		Bt1G3231	Bt1G3232		Bt1G3233	Bt1G3234 1		1 CC7CD110	Bt1G3236	Bt1G3237	Bt1G3238	Bt1G3239	Bt1G3240	Bt1G3241	Bt1G3242	Bt1G3243	Bt1G3244	Bt1G3245	
Position	5895-5024	6479-5898		7882-6482	9031-8111		10014-9137	10812-10114		120011-10021	1-865	1241-1972	1985-2971	3866-5392	5982-6968	6997-7827	7857-8732	9088-9786	330-1	1068-352	
NCBI gi	g129533	g129521		g129527	g585031		g586894	g586893		8202129	g2226185	g2226242	g2226243	g2226245	g4323056	g118163	g118165	g2462099	g2633746	g2633745	
aat_ nap Score	710	789		1384	1268		923	892	} }.	1505	489	756	1264	1559	1187	733	725	1142	496	818	
BlastP Score	747	793		1232	1132		906	898		1234	510	761	1274	1547	1122	580	707	1081	496	833	
BlastP- Prob	5.30E-74	7.10E-79		2.10E-125	8.40E-115		7.50E-91	5.30E-90		1.50E-125	6.90E-49	1.70E-75	7.60E-130	8.90E-159	9.70E-114	2.60E-56	9.20E-70	2.10E-109	2.10E-47	4.10E-83	
% % Ident Cvrg	49	77	•	58	8		62	70	}	7	36	59	72	58	68	51.	48	95	8	2	
Cvrg	100 /	99]	3 ra b	100	000		100	100		2	86 (100	100 (100	100	100	100	66	100	
NCBI gi description	[Bacillus subtilis] 4-AMINO-4-DEOXYCHORISMATE LYASE (ADC	LYASE) [Bacillus subtilis] PARA-AMINOBENZOATE SYNTHASE GLUTAMINE	AMIDOTRANSFERASE COMPONENT II (ADC SYNTHASE) / ANTHRANILATE SYNTHASE	COMPONENT II [Bacillus subtilis] PARA-AMINOBENZOATE SYNTHASE COMPONENT I	(ADC SYNTHASE) [Bacillus subtilis] CYSTEINE SYNTHASE (O-ACETYLSERINE	SULFHYDRYLASE) (O-ACETYLSERINE (THIOL)- LYASE) (CSASE) (SUPEROXIDE-INDUCIBLE PROTEIN	11) (SOI11) [Bacillus subtilis] HYPOTHETICAL 31.8 KD PROTEIN IN FTSH-CYSK	INTERGENIC REGION [Bacillus subtilis] HYPOTHETICAL 26.2 KD PROTEIN IN FTSH-CYSK	INTERGENIC REGION [Bacillus subtilis]	subtilis]	(Y14081) hypothetical protein [Bacillus subtilis]	(Y14083) hypothetical protein [Bacillus subtilis]	(Y14083) hypothetical protein [Bacillus subtilis]	(Y14083) hypothetical protein [Bacillus subtilis]	(AF098509) putative sulfur-binding protein [Enterobacter cloacae]	SULFATE TRANSPORT SYSTEM PERMEASE PROTEIN CYST [Escherichia coli]	SULFATE TRANSPORT SYSTEM PERMEASE PROTEIN CYSW [Escherichia coli]	(Y10908) sulfate transport ATP-binding protein [Bacillus	cereus] (Z99111) similar to hypothetical proteins [Bacillus subtilis]	(Z99111) similar to coenzyme PQQ synthesis [Bacillus	Subtitude

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1352 1352	1352	1351	1351	1351	1351	1351	1351	1351		1351	1351	1351	1351	1351	1350	1330				1350		1350	1350	1350	1350	S E S	
1352 Bt1Gc1699 1352 Bt1Gc1699	Bt1Gc1699	Bt1Gc1698 Bt1G3268	Bt1'Gc1698	Bt1Gc1698	Bt1Gc1698	Bt1Gc1698	Bt1Gc1698	Bt1Gc1698		Bt1Gc1698	Bt1Gc1698	Bt1Gc1698	Bt1Gc1698	Bt1Gc1698	Bt1Gc1696	96100119	Bt1Gc1696	Bt1Gc1696	Bt1Gc1696	Bt1Gc1696		Br1 Gc1696	Bt1Gc1696	Bt1Gc1696	Bt1Gc1696	Contig Id	
Bt1G3270 Bt1G3271	Bt1G3269		Bt1G3267	Bt1G3266	Bt1G3265	Bt1G3264	Bt1G3263	Bt1G3262		Bt1G3261	Bt1G3260	Bt1G3259	Bt1G3258	Bt1G3257	Bt1G3256	CC7CD119	Bt1G3254	Bt1G3253	Bt1G3252	Bt1G3251		B+1G3250	Bt1G3249	Bt1G3248	Bt1G3247	Gene Id	
1339-2004 2030-3715	324-1346	20161-18279	12557-14497	9876-12539	9153-9692	8595-9023	7062-8588	6269-6628		4027-5787	3023-3793	2554-2811	1610-2401	1-1444	17461-16819	13393-14213	15206-12892	13141-9996	11764-11144	9441-10271		7130-8650	6937-6503	6361-4667	2211-1555	Position	
g2635770 g2635769	g2635771	g1731026	7 g1709188	g2634076	g116957	g2634074	g2634073	g2462119	. ,	g2129427	g2062103	g1174408	g2634069	g2634068	g1763079	80000009				g1176306	9	02633737	g2633738	g3323354	g2633743	NCBI gi	
773 793	1170	1650	1927	2799	562	484	2124	609		597	583	382	1021	2120	596	230	421	335	372	264		014	381	1325	862	nap Score	
479 787	1136	1685	1248	2761	500	485	2124	609		709	609	312	1021	1851	642 242	249	466	437	400	344		96	358	1257	871	BlastP Score	
1.30E-45 3.10E-78	3.20E-115	2.10E-173	8.60E-188	2.00E-287	7.90E-48	3.10E-46	6.40E-220	2.20E-59		1.60E-83	2.20E-59	6.60E-28	4.90E-103	5.40E-191	7.10E-63	3.10E-21	1.60E-43	3.70E-41	3.10E-37	2.70E-31		3 30F-97	8.80E-33	5.20E-138	3.80E-87	BlastP- Prob	
68	66	51	60	·65	58	66	78	98		34	.47	92	72	88	59	29	36	39	39	30		37	54	45	71	% Ident	
100	100	98	100	100	B	99	100	99		100	100	99	100	93	58	. 6	39	57	100	100		3 .	99	100	100	Cyrg	
[Bacillus subtilis] (Z99120) similar to hypothetical proteins [Bacillus subtilis] (Z99120) similar to hypothetical proteins [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis] (Z99120) similar to ABC transporter (ATP-binding protein)	subtilis] HYPOTHETICAL 73.2 KD PROTEIN IN SODA-COMGA	DNA MISMATCH REPAIR PROTEIN MUTL [Bacillus	(Z99112) mutS [Bacillus subtilis]	SPORE COAT PROTEIN E [Bacillus subtilis]	(Z99112) similar to hypothetical proteins [Bacillus subtilis]	(Z99112) similar to hypothetical proteins [Bacillus subtilis]	(Y11140) bc48a [Bacillus cereus]	- Sulfolobus acidocaldarius [Sulfolobus sp. 7]	2-oxoacid ferredoxin oxidoffeductase (EC 1) alpha chain	(X93169) orfD [Listeria monocytogenes]	STAGE V SPORULATION PROTEIN S [Bacillus subtilis]	(Z99112) similar to hypothetical proteins [Bacillus subtilis]	(Z99112) similar to hypothetical proteins [Bacillus subtilis]	(U69493) PhnW [Salmonella typhimurium]	(299111) transcriptional regulator (Deok family) [Bacillus	INTERNALIN A PRECURSOR []	(AF065404) pXO1-54 [Bacillus anthracis]	(Z99111) similar to hypothetical proteins [Bacillus subtilis]	HYPOTHETICAL PROTEIN HI0755 []	[Bacillus subtilis]	[Bacillus subtilis]	pallidum (Z99111) similar to transcriptional regulator (MarR family)	(AE001270) oligoendopeptidase F, putative [Treponema	influenzae Rd] (Z99111) similar to hypothetical proteins [Bacillus subtilis]	NCBI gi description	-

1355]	1354 I	1354 1	1334 1		1354 1	1354 I	1354 I	1354 I	· 1354 I	1354 I	1354 I	1354 I	1353 I	-	1353 I		٠.•	1353 I	1000		1353 I	1353 I		- 1353 I	1352 I	1352 I	1352 I	1352 I	·1352 I	1352 I	SEQ NO	
Bt1Gc1695	Bt1Gc1692	Bt1Gc1692	811001092	011001092	B+10-1602	Bt1Gc1692	Bt1Gc1692	Bt1Gc1692	Bt1Gc1692	Bt1Gc1692	Bt1Gc1692	Bt1Gc1692	Bt1Gc1700	٠	Bt1Gc1700			Bt1Gc1700	P(10C1/00	3+10-1700	Bt1Gc1700	Bt1Gc1700		Bt1Gc1700	Bt1Gc1699	Bt1Gc1699	Bt1Gc1699	Bt1Gc1699	Bt1Gc1699	Bt1Gc1699	Contig Id	
Bt1G3296	Bt1G3295	Bt1G3294	B[103293			Bt1G3291	Bt1G3290	Bt1G3289	Bt1G3288	Bt1G3287	Bt1G3286	Bt1G3285	Bt1G3284		Bt1G3283			Bt1G3282	D(1/02/01	D+1C2201	Bt1G3280	Bt1G3279		Bt1G3278	Bt1G3277	Bt1G3276	Bt1G3275	Bt1G3274	Bt1G3273	Bt1G3272	Gene Id	
1376-2221	10426-8711	8721-7896	C00/-/18/	7017 7000	7815-7189	5349-4059	4741-3576	4747-2525	3965-3600	3432-2323	2067-1354	1640-260	12485-11529		10106-8187			8218-7079	/0/0-0000	7076 6060	5926-4478	4477-2081		1-214	11285-11587	9816-11195	7740-9134	6050-7267	4758-6047	3957-4739	Position	
g2635736	g2293146	g2293147	82293148	2223149	-2203140	g2495742	g1002992	g3426013	g1149666	g2293150	g4481749	g282286	9 g126054		g98191	•.		g729582	10011078	2011061	_. g2811062	g1944414	-	g136144	7 g2635733	g2635734	ģ2635763	g2635765	g2635766	g3183561	NCBI gi	
932	1355	568	1233	1203	1262	390	318	330	367	1034	539	427	1137		2370			1475	772	3.	1589	2976		316	345	1027	2232	1726	1445	1221	nap Score	
781	1305	565	1118	119/	1107	418	460	510	371	1051	547	514	1109		2269			1484	723	3	1608	2926		340	350	1019	2136	1674	1339	1221	BlastP Score	
1.30E-77	3.90E-133	1.00E-54	2.60E-113	1.10E-121	1 105 131	3.90E-39	1.40E-43	6.90E-49	3.70E-34	3.20E-106	8.30E-53	2.60E-49	2.30E-112		2.80E-235			4.20E-152	1.200-92	305	3.10E-165	6.60E-305		7.10E-31	6.20E-32	7.90E-103	3.40E-221	3.10E-172	9.80E-137	3.10E-124	BlastP- Prob	
56	46	43	%	21		. 40	34		60	54	45	.34	66		65		,	71	. 0/	ĵ	62	71		94	70	45	91	78	.67	90	% Ident	
100	99	100	. 100	100	3. 3.	100	100	94	99	100	100	100	100		100	•		100	100	3	100	100		15	99	100	100	100	100	100	Cvrg	
(Ź99120) yunF [Bacillus subtilis]	(AF008220) SHCHC synthase [Bacillus subtilis]	(AF008220) YtxM [Bacillus subtilis]	(AF008220) dinydroxynaphthoate synthase [Bacillus subtilis]	(Arovozzo) Con-Con synthase [Dacinus suonns]	jannaschii]	HYPOTHETICAL PROTEIN MJ0014 [Methanococcus	(U34772) ORF375 [Dichelobacter nodosus]	(AB016803) transposase [Deinococcus radiodurans]	(X86498) IS1136 DNA [Clostridium perfringens]	(AF008220) YtfD [Bacillus subtilis]	(AF007865) BacR [Bacillus licheniformis]	VanS - Enterococcus faecium []	L-LACTATE DEHYDROGENASE X []	stearothermophilus [Bacillus stearothermophilus]	1,4-alpha-glucan branching enzyme (EC 2.4.1.18) - Bacillus	PYROPHOSPHORYLASE) [Bacillus subtilis]	(ADP-GLUCOSE SYNTHASE) (ADP-GLUCOSE	STEATOTH STEAT STE	OLICOGEN BIOSINI DESIS FROI EIN GLOD (Bacillus	GLYCOGEN) SYNTHASE) [Bacillus stearothermophilus]	STARCH (BACTERIAL GLYCOGEN SYNTHASE (STARCH (BACTERIAL	(D87026) glycogen phosphorylase [Bacillus	ELEMENT IS231C [Bacillus thuringiensis]	TRANSPOSASE FOR INSERTION SEQUENCE	(Z99120) yunC [Bacillus subtilis]	(Z99120) similar to hypothetical proteins [Bacillus subtilis]	(Z99120) similar to hypothetical proteins [Bacillus subtilis]	(Z99120) similar to NifS protein homolog [Bacillus subtilis]	(Z99120) similar to hypothetical proteins [Bacillus subtilis]	VEGETATIVE PROTEIN 296 (VEG296) [Bacillus subtilis]	NCBI gi description	

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	Bt1Gc1702 Bt1Gc1702	Bt1Gc1702	Bt1Gc1702	Bt1Gc1702	Bt1Gc1702	Bt1Gc1702	Bt1Gc1695	Bt1Gc1695	Bt1Gc1695	Bt1Gc1695		Bt1Gc1695	Bt1Gc1695	Bt1Gc1695	••		Bt1Gc1695	Bt1Gc1695	Bt1Gc1695	Bt1Gc1695	Bt1Gc1695		Bt1Gc1695		Bt1Gc1695	Contig Id
	Bt1G3316 Bt1G3317	Bt1G3315	Bt1G3314	Bt1G3313	Bt1G3312	Bt1G3311	Bt1G3310	Bt1G3309	Bt1G3308	Bt1G3307		Bt1G3306	Bt1G3305	Bt1G3304	,		Bt1G3303	Bt1G3302	Bt1G3301	Bt1G3300	Bt1G3299		Bt1G3298	•	Bt1G3297	Gene Id
	12624-11719 g2632986 14612-13188 g2589196	10660-9353	9231-7852	7838-6396	4121-5350	438-1631	19101-19820	17744-18800	16009-17641	15353-15896		14528-15214	13708-14241	13115-13702			11989-13124	10983-11882	10320-10859	9777-9019	9033-7195		6227-5523		5772-3403	Position
	9 g2632986 8 g2589196	10660-9353 g1805459	g1072970	g120777	g2507254	g2636487	0 g466194	0 g466193	1 g466192	6 g466191				2 g466188			4 g461914	2 g2619018	9 g466187	g2633166	g2804528	(£2126871		g2500765	NCBI gi
	1023 2081	1472	797	1413	1410	775	965	1286	1836	427		826	578	705®			851	762	332	430	1551		923		301	nap Score
	1038 2090	1374	788	1330	1415	675	844	1293	1807	462			614	654			796	752	366	415	1492		941		395	BlastP Score
	7.70E-105 2.60E-216	1.90E-140	2.40E-78	8.80E-136	8.60E-145	2.30E-66	2.80E-84	7.30E-132	2.50E-186	8.40E-44		6.60E-83	6.60E-60	3.80E-64			3.40E-79	1.60E-74	1.20E-33	8.00E-39	6.00E-153		1.50E-94		1.10E-36	BlastP- Prob
	865	63	39	56	2	42	77	68	63	46		69	66	67			48	52	41	38	51		78		29	% Ident
	100	100	100	100	100	100	100	100	100	99		100	99	99			100	100	99	100	100		100		67	% Cyrg
•	GabT of E. coli [Bacillus subtilis] (Z99107) similar to hypothetical proteins [Bacillus subtilis] (AF008553) Glu-tRNAGIn amidotransferase subunit B	autotrophicus] (D50453) homologue of 4-aminobutyrate aminotransferase	(NADP+) (SSDH) [Escherichia coli] dhlR protein - Xanthobacter autotrophicus [Xanthobacter	SUCCINATE-SEMIALDEHYDE DEHYDROGENASE	AMINOPEPTIDASE AMPS [Bacillus subtilis]	(Z99124) pyrimidine-nucleoside transport protein [Bacillus subtilis]	TRANSCRIPTIONAL REGULATORY PROTEIN RESD [Bacillus subtilis]	RESC PROTEIN [Bacillus subtilis]	RESB PROTEIN [Bacillus subtilis]	RESA PROTEIN [Bacillus subtilis]	SYNTHASE B (PSEUDOURIDYLATE SYNTHASE) (URACIL HYDROLYASE) [Bacillus subtilis]	RIBOSOMAL LARGE SUBUNIT PSEUDOURIDINE	SPORE MATURATION PROTEIN B [Bacillus subtilis]	SPORE MATURATION PROTEIN A [Bacillus subtilis]	PEPTIDASE) (DD-CARBOXYPEPTIDASE) (PBP-5*) [Bacillus subtilis]	ALANYL-D-ALANINE CARBOXYPEPTIDASE) (DD-	PENICILLIN-BINDING PROTEIN 5* PRECURSOR (D-	INTERGENIC REGION (ORFX9) [Bacillus subtilis] (AF027868) superoxide dismutase [Bacillus subtilis]	HYPOTHETICAL 20.4 KD PROTEIN IN RIBT-DACB	(Z99108) yfiW [Bacillus subtilis]	(D85082) YfiX [Bacillus subtilis]	stearothermophilus [Bacillus stearothermophilus]	purine nucleoside phosphorylase (EC 2.7) II - Bacillus	RESISTANCE PROTEIN VANSB) (VANCOMYCIN	SENSOR PROTEIN VANSB (VANCOMYCIN B-TYPE	NCBI gi description
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	Bt1Gc1704	Bt1Gc1704	Bt1Gc1,704	Bt1Gc1704	Bt1Gc1704		Bt1Gc1704	•		Bt1Gc1704	Bt1Gc1701	Bt1Gc1701	Bt1Gc1701		Bt1Gc1701	Bt1Gc1701	į	Bt1Gc1701	Bt1Gc1701	Bt1Gc1701		Bt1Gc1701	Bt1Gc1701	Bt1Gc1701	Bt1Gc1701	Bt1Gc1701		Bt1Gc1701	Bt1Gc1702		Contig Id	
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	Bt1G3339	Bt1G3338	Bt1G3337	Bt1G3336	Bt1G3335		Bt1G3334			Bt1G3333	Bt1G3332	Bt1G3331	Bt1G3330	•	Bt1G3329	Bt1G3328		Bt1G3327	Bt1G3326	Bt1G3325	-	Bt1G3324	Bt1G3323	Bt1G3322	Bt1G3321	Bt1G3320		Bt1G3319	Bt1G3318		Gene Id	
	6807-10716	7439-7915	5962-7146	5107-5910	2984-4291		1146-2990			438-1139	16972-16412	16936-16415	12396-14729		11096-11980 g131027	10368-10030		9467-11001	8805-9158	8194-8628		7625-6885	6895-5169	5664-4954	3792-3316	3329-1881		1680-1026	16084-14630 g2632982		Position	
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	g2633724	g1064795	g1731364	g1064810	g2636586		g1064813			g586808	g548832	g3024615	g2635858	•	131027	g2226121		g132549	g2226119	g3023940	•	g1706577	g1706578	g1706579	g1881269	g1881270		g322156	632982	·	NCBI gi	
	353	640	929	979	264		1694		:'	957	157	156	1830		564	210		615	214	524		892	609	594	359	530		384	2025		nap Score	
	436	640	929	983	429		1649			864	172	208	1833		284	226		659	215	491	•	851	610	497	363	655		388	1875		BlastP Score	
	1.70E-43	1.20E-62	2.70E-93	5.20E-99	2.60E-40		1.40E-169			2.10E-86	4.50E-13	6.90E-17	4.40E-189		5.00E-33	8.60E-19		1.10E-64	1.30E-17	7.10E-47		5.00E-85	1.70E-59	1.60E-47	2.60E-33	3.00E-64		5.80E-36	1.60E-193		BlastP- Prob	
	43	52	93	- 99	4		169			-86	13	-17	189		-33	-19		2	-17	47		- 85	-59	47	' 33	2		-36	193		•	-
	36	77	4 %	.71	26		55			79	26	25	46		45	36		60	31	66		86	36	49	43	32		42	.80		% % Ident Cvrg	
	35	99	100	100	100		100			100	99	99	100		100	99		100	99	. 99		100	100	1 00	99	100		100	100		%gr	
[Bacıllus subtilis]		INTERGENIC REGION [Bacillus subtilis] (D78193) yydA [Bacillus subtilis]		(D78193) yycJ [Bacillus subtilis]	(Z99124) yycH [Bacillus subtilis]	subtilis]		[Bacillus subtilis]	PROTEIN IN ROCR-PURA INTERGENIC REGION	HYPO'		RNA POLYMERASE SIGMA FACTOR SIGV [Bacillus	(Z99121) yvgS [Bacillus subtilis]	[Bacillus subtilis]		(Y14077) Hypothetical protein [Bacillus subtilis]	HPR [Bacillus subtilis]	PROTEASE PRODUCTION REGULATORY PROTEIN	(Y14077) Hypothetical protein [Bacillus subtilis]	HIT PR	ECSA [Ba	_		PROTE	(AB00	(AB00	megaterium]	Bacillus subfilis hypothetical protein 1 - Bacillus megaterium Bacillus		[Bacillus subtilis]		
ubtilis	simila	yydA YydA	ETIC/	χyc	уусН		homo	ubtilis	IN R	omonas aeruginosaj THETICAL 27.2 KD SENSORY TRANSDUCTION	YME	YME	yvgS	ubtilis	EXP	Hypot	illus s	E PR	Hypot	EIN	[Bacillus subtilis]	ETR	ECSI	ECS(1488) FUNCTION UNKNOWN. [Bacillus subtilis]	1488) FUNCTION UNKNOWN. [Bacillus subtilis]	브	al pro	alterna	ubtilis	•.	
بق	rtot	Ba	AL P	Baci	[Bac		logou	ت	OCR.	erugi	RASI	RASI	Bac	<u>~</u>	ORT	hetic	ubtil	ndo	hetic	Вас	subt	ANSI	뗪	ਯੂ	NCT	NCT		tein 1	ite ge	ٽ	Z	
	WO-C	illus	ROT	llus s	illus		s to		-PUR	nosa 7.2 K	SIS 3	E SIC	illus		PRO	al pro	<u>is</u>	CTIC	al pro	illus	ilis]	PORT	acillu	acillu	Ö	ÖN		- Ba	ne na		NCBI gi description	
٠.	ompo	Bac	EASI	ubtil	subti		sp:PF		A	D SE	ΔMA	ΜĂ	subti		TEIP	otein		NR	otein	subtil		ER.	s sub	s sub	SN	Ş		cillu	me:		gi de	
	nent	llus s	Z	<u>ıs</u>	lis]		Ŗ.		TER	NSO	·H F/	 FAC	<u>is</u>		PRS	[Bac		EGU	[Bac	īs		ATP-	tilis]	tilis]	NON	NOV		s meg	yedB		scrip	
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	or his	· <u> s</u>	. R-PU				US		ICR	[RA]	OR (S	SIG			SEC(subti		ORY	subti			DINC			Baci	Baci		E E	ilar t			•
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100 HYPOTHETICAL ABC TRANSPORTER PERMEASE	42 1	2.50E-24	236	447	g401471	1361 Bt1Gc1708 Bt1G3376 13670-13020 g401471	Bt1G3376	1 Bt1Gc1708	136
subthis] 100 (Z99110) alternate gene name: yidA; similar to 3-oxoacyl-	54 1	1.40E-71	724	653	g2633552	12205-12978	Bt1G3375	1361 Bt1Gc1708	136
[Bacillus subrins] 100 (Z99120) similar to acylaminoacyl-peptidase [Bacillus	26 1	6.30E-59	411	420	g2635720	8601-10517	Bt1G3374	1361 Bt1Gc1708	136
(GFAT) (L-GLUTAMINE-D-FRUCTOSE-6-PHOSPHATE AMIDOTRANSFERASE) (GLUCOSAMINE-6-PH					٠				
AMINOTRANSFERASE (ISOMERIZING) (HEXOSEPHOSPHATE AMINOTRANSFERASE) (D-	-								
[Bacillus subtilis] 100 GLUCOSAMINEFRUCTOSE-6-PHOSPHATE	74 1	6.30E-229	2209	2277	g1169919	6511-8310	Bt1G3373	1 Bti/Gc1708	1361
100 (Z99104) similar to phosphoglucomutase (glycolysis)	79 1	2.90E-185	1797	1857	g2632444	4678-6021	Bt1G3372	1 Bt1Gc1708	1361
100 (Y15477) hypothetical protein [Staphylococcus aureus]	29 1	7.40E-36	387	304	g3892894	3207-4163	Bt1G3371	1 Bt1Gc1708	1361
100 (AB002150) YbbP [Bacillus subtilis]	68 1	1.80E-98	978	978	g1944009	2384-3202	Bt1G3370	1 Bt1Gc1708	1361
100 ARGINASE [Bacillus caldovelox]	74 1	1.50E-115	1139	1118	g1703387	1245-2132	Bt1G3369	1 Bt1Gc1708	1361
					(
		1.10E-91		958		1-1031			1361
	31 1	2.90E-34	•	297		10124-10903	Bt1G3367	Bt1Gc1705	1360
99 (Z99120) yuzD [Bacillus subtilis]	47	1.30E-24	281	257	g2635718	9733-10062	Bt1G3366	Bt1Gc1705	1360
subtilis]	,		,						
99 (Z99120) similar to NifU protein homolog [Bacillus	68	2.90E-34	372	347	g2635719	9641-9313	Bt1G3365	1360 Bt1Gc1705	136
100 (Z99120) yutH [Bacillus subtilis]	41 1	3.80E-71	720	705	g2635724	7834-8850	Bt1G3364	0 Bt1Gc1705	1360
[Bacillus subtilis]									
		6.00E-50	520	582	g2635725	7725-7228	Bt1G3363	1360 Bt1Gc1705	136
99 (US9239) orf3 [Serratia marcescens]	37	1.80E-25	289	180	g1695869	6376-4777	Bt1G3362) Bt1Gc1705	1360
				. ;	0				. ;
92 PUTATIVE NAD(P)H OXIDOREDUCTASE HI1544	37	7.70E-34	368	343	g1175888	1631-1036	Bt1G3361	1360 Bt1Gc1705	136
100 (Z99120) similar to N-acetyl-glucosamine catabolism	69 1	8.00E-87	868	907	g2635726	14-781	Bt1G3360) Bt1Gc1705	1360
								. •	
[Bacillus subtilis] 56 HISTIDINOL DEHYDROGENASE (HDH) [Bacillus	57	4.30E-63	64	666	g3122219	20555-19836	Bt1G3359	9 Bt1Gc1703	1359
99 (AF017113) imidazoleglycerol-phosphate dehydratase	51	2.70E-47	495	495	g2618867	19836-19255	Bt1G3358	9 Bt1Gc1703	1359
R NCBI gi description	% % Ident Cvrg	Prob	Score Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	S E
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(AF053927) probable spore germination protein E [Bacillus cereus]	. 99	100	2.30E-64	656	656	g2984722	4393-4776	Bt1G3417	5 Bt1Gc1712	1365
(AF053927) probable spore germination protein D [Bacillus	98	100	9.40E-29	320	320	g2984721	4183-4374	Bt1G3416	5 Bt1Gc1712	1365
(AF053927) probable spore germination protein C [Bacillus	100	99	2.70E-95	948	1047	g2984720	3562-4173	Bt1G3415	5 Bt1Gc1712	1365
(AF053927) probable spore germination protein B [Bacillus	99	99	5.10E-21	247	327	g2984719	3288-3491	Bt1G3414	5 Bt1Gc1712	1365
(AF053927) probably spore germination protein A [Bacillus	. 99	99.	7.70E-34	368	368	g2984718	3052-3270	Bt1G3413	5 Bt1Gc1712	1365
(AF053927) YisI homolog [Bacillus cereus]	98 (93	1.80E-25	289	289	g2984717	2776-2952	Bt1G3412	5 Bt1Gc1712	1365
(Y09476) YisK [Bacillus subtilis]	100	54	4.80E-80	804	869	g2145376	2529-1633	Bt1G3411	5 Bt1Gc1712	.1365
ORNITHINE AMINOTRANSFERASE (ORNITHINE-	100	75	2.30E-167	1628	1617	g730217	1-1203	Bt1G3410	btlGc1/12	1365
(U94743) hemolysin II [Bacillus cereus]		39	3.60E-52	541	486		13556-15849	Bt1G3409		1364
(D90770) ORF_ID:o260#11; similar to [Escherichia coli]	100 (42	8.40E-60	613	537		12577-11656	Bt1G3408	1 Bt1Gc1709	1364
CHROMOSOME X [Caenorhabditis elegans]	_					,				
mermoautotropnicum] HYPOTHETICAL 69.0 KD PROTEIN F40B5.2 IN	55 1	27	1.20E-14	201	137	g2496971	12821-8950	Bt1G3407	Bt1Gc1709	1364
(AE000886) conserved protein [Methanobacterium	100	29	1.20E-28	319	197	g2622286	11508-10394	Bt1G3406	1 Bt1Gc1709	1364
(D88802) ydhJ [Bacillus subtilis]	100 (55	6.40E-101	1001	999	g1945094	10265-9291		Bt1Gc1709	1364
coelicolor]	•									
(AJ010320) RNA polymerase sigma factor [Streptomyces	81 (28	1.90E-14	185	86	g3688548	7678-8180	Bt1G3404	1 Bt1Gc1709	1364
(Y14082) hypothetical protein [Bacillus subtilis]	99 (40	2.50E-28	316	244	g2226208	7641-8124	Bt1G3403	# Bt1Gc1709	1364
lactis]	_				٠.	•				
(AE001272) positive transcriptional activator [Lactococcus	100.	36	1.40E-43	460	331	g3582223	7419-6588	Bt1G3402	1 Bt1Gc1709	1364
(277120) suntia to two-component sensor manume surase	5	0	0.702-100	1000	1000	0100009	1101-1017	1010111	, 511001707	
[Bacillus subtilis]		٠ •	0 20E 100	1066	1068	~7625010	1131-5917			136
(Z99120) similar to two-component response regulator	71 (55	1.20E-69	706	650	g2635819	3338-4115	Bt1G3400	# Bt1Gc1709	1364
(Y14083) hypothetical protein [Bacillus subtilis]		51	1.20E-117	1159	1156	g2226247	1489-212	Bt1G3399	1 Bt1Gc1709	1364
burgdorferi]	~				. :					
(AE001172) NADH oxidase, water-forming (nox) [Borrelia		39	4.00E-92	918	880	g2688656	12532-13866	_		1363
[Lactococcus lactis] (Z82015) vukJ [Bacillus subtilis]	9 [6	5.90E-27	303	301	g1665854	8333-8771	Bt1G3397	Bt1Gc1711	1363
					Score	٠		-	-	NO
NCBI gi description	Cyrg	% Ident	BlastP- Prob	BlastP Score	aat nap	NCBI gi	Position	Gene Id	Contig Id	SEQ
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		1366 Bt1Gc1713	1366 Bt1Gc1713	1366 Bt1Gc1713	1366 Bt1Gc1713		1366 Bt1Gc1713	٠.	1366 Bt1Gc1713	1366 Bt1Gc1713		1366 Bt1Gc1713	1366 Bt1Gc1713	1366 Bt1Gc1713	1366 Bt1Gc1713		1366 Bt1Gc1713		1366 Bt1Gc1713		1366 Bt1Gc1713	1366 Bt1Gc1713	1366 Bt1Gc1713	1365 Bt1Gc1712	1365 Bt1Gc1712	1365 Bt1Gc1712	1365 Bt1Gc1712	NO	SEQ ID Contig Id
•		1713	1713	1713	1713		1713		1713	1713		1713	1713	1713	1713		1713		1713		1713	1713	1713	1712	1712	1712	1712		g Id
		Bt1G3437	Bt1G3436	Bt1G3435	Bt1G3434		Bt1G3433		Bt1G3432	Bt1G3431	_	Bt1G3430	Bt1G3429	Bt1G3428	Bt1G3427		Bt1G3426		Bt1G3425		Bt1G3424	Bt1G3423	Bt1G3422	Bt1G3421	Bt1G3420	Bt1G3419	Bt1G3418		Gene Id
•		14693-13890 g3122350	13825-13178 g2618860	13132-12619 g2618861	12480-10972		10883-9936		9755-8952	8734-8261		8252-7369	7406-6406	6320-5373	5346-5092		4189-4779		3091-3792		2490-2843	1474-2490	1021-1470	13305-12841	12712-9191	9194-5484	4822-5034		Position
) g3122350	8 g2618860	g2618861	2 g1945645		g3123300		g2688692	g1945649	•	g2811049	g2635989	g1945652	g1945653		g3287962	٠.	g1881360		g730786	g730785	g730784	g729934	g113346	g113345	g2984723	•	NCBI gi
		972	717	709	692		1411		182	578		1026	1063	1320	293		877	`.	348		335	805	459	543	2954	3258	366	₹₽	aat_
		807	662	725	745		1411		148	592		1049	1060	1320	259	•	883	. -	370		385	831	468	546	2954 2990 1.1e-311	3322	292	26016	BlastP
		2.30E-80	5.40E-65	1.10E-71	8.60E-74		2.30E-144		4.00E-10	1.40E-57		5.30E-106	3.60E-107	1.00E-134	2.70E-22		2.00E-88		4.70E-34		1.20E-35	6.60E-83	1.90E-44	1.10E-52	l.1e-311	0	8.70E-26	1.00	BlastP-
		67	63	78	36		84		29	66		68.	2	80	67		&		34		57	4	57	63	50	54	100	100110	Ident
		100	100	99	100		100		100	.99		100	100	100	99		.99		100		37	100	99	85	100	100	99	61.6	3 %
	TRANSFERASE (SPORE GERMINATION PROTEIN GERF) [Bacillus subtilis]	PROL	_	_	_	PROTEIN 35) (GSP35) [Bacillus subtilis]		[Borrelia burgdorferi]	_	(Z940	INTERGENIC REGION [Bacillus subtilis]	_			(Z94043) hypothetical protein [Bacillus subtilis]	PROTEASE) (PROTEASE TI) [Bacillus subtilis]	ATP-I	THIS ENTRY (YDFR_BACSU). [Bacillus subtilis]		subtilis]			I) [Ba STAC	subtilis] SIGNAL PEPTIDASE I (SPASE I) (LEADER PEPTIDASE					NCBI gi description

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(PHOSPHOHEXOKINASE) [Bacillus stearothermophilus] ACETYL-COENZYME A CARBOXYLASE CARBOXYL	2 100	1 72	4.30E-111	1097	1245	g3121 73 6	3947-2973	Bt1G3458	Bt1Gc1706	1368
6-PHOSPHOFRUCTOKINASE			1.00E-134		1422	g125125	S	Bt1G3457		1368
(AJ222587) YkuP protein [Bacillus subtilis] DVB1IVATE VINACE (DV) [Bacillus chearathermorphilus]	100	60 E 4 E	4.50E-29	323 1684	350	g2632237	19817-20665	Bt1G3455	Bt1Gc1714	1367
(AMINOACYLASE) [Bacillus stearothermophilus]	36		9.00		Ş	000111				į
N-ACVI-I-AMINO ACID AMIDOHYDROI ACE		5 2	8 605-90	80%	803	o584740	18245_19347 o584749	Rt1G3454	7 Rt1Gc1714	1367
ALANINE-D-ALANYL CARRIER PROTEIN LIGASE)						٠.				
D-ALANINE-ACTIVATING ENZYME (DAE) (D-	6 100	19 56	1.50E-149	1460	1435	g729340	17751-16240 g729340	Bt1G3453	/ Bt1Gc1714	1367
DLTB PROTEIN [Bacillus subtilis]	9 100	9 59	2.70E-109	1080	1164	g729341	16240-15065	Bt1G3452	1 Bt1Gc1714	1367
(D86240) D-alanyl carrier protein [Staphylococcus aureus]	9 99	5 69	1.80E-25	289	288	g1405337	14993-14760	Bt1G3451	7 Bt1Gc1714	1367
PROTEIN DLTD PRECURSOR [Bacillus subtilis]	8 100	3 48	6.20E-103	1020	1013	g729343	14760-13573	Bt1G3450	1 Bt1Gc1714	.1367
(U93875) 2-nitropropane dioxygenase [Bacillus subtilis]	0 100	_	4.80E-57	587	588	g1934639	12481-13524	Bt1G3449	1 Bt1Gc1714	1367
(AJ002571) YkfA [Bacillus subtilis]	3 100	7 43	3.40E-47	494	442	g2632018	12125-11440 g2632018	Bt1G3448	1 Bt1Gc1714	1367
(U93876) hypothetical protein YrdC [Bacillus subtilis]	2 99	l6 52	6.40E-46	482	470	g1934645	10372-10935	Bt1G3447	7 Bt1Gc1714	1367
REGULATOR. [Bacillus subtilis]										
REGULATOR, SIMILAR TO LEU RESPONSIVE						٠			,	
(AB001488) PROBABLE TRANSCRIPTIONAL	4 77	ō 4	4.30E-40	427	357	g1881235	11088-9393	Bt1G3446	Bt1Gc1714	1367
DROTER! [Resiling enterlial	100	5	0.7015-00	777	9	81001237	0777-7005	P(10)440		į
[Helicobacter pylori 26695]		-	3 00E	303	707	~1001750	60/17-7863	B+1/C2///\$	B+1C-1714	1367
(AE000627) ABC transporter, ATP-binding protein (yhcG)	2 100	3 42	1.40E-43	460	495	g2314379	5461-6147	Bt1G3444	Bt1Gc1714	1367
subtilis]										
IN PRKA-CSPB INTERGENIC REGION [Bacillus								•		
[Bacillus subtilis] HYPOTHETICAL TRANSCRIPTIONAL REGULATOR	2 99	3 42	1.10E-23	272	266	g1724000	5078-5440	Bt1G3443	⁷ Bt1Gc1714 Bt1G3443	1367
BETA CHAIN (RIBONUCLEOTIDE REDUCTASE)										
[Staphylococcus aureus] RIBONUCLEOSIDE-DIPHOSPHATE REDUCTASE	7 100	15 57	1.00E-95	952	892	g1710395	3443-4873	Bt1G3442	Bt1Gc1714	1367
(AJ133495) ribonucelotide reductase major subunit	4 100	3 64	1.40E-233	2253	2358	g4490609	1146-3297	Bt1G3441	Bt1Gc1714	1367
EXCINUCLEASE ABC SUBUNIT A [Bacillus subtilis]	2 44	9 82	2.10E-189	1836	1815	g3915204	18080-16814	Bt1G3440		. 1366
(AF017113) YvID [Bacillus subtilis]	1 99	6 51	3.00E-16	202	311	g2618847	16192-15821 g2618847	Bt1G3439	Bt1Gc1713	1366
HPR(SER) KINASE [Bacillus subtilis]	5 100	7. 75	5.00E-117	1153	1211	g3122218	15652-14723	Bt1G3438		1366
NCBI gi description	t Cyrg	% Ident	BlastP- Prob	BlastP Score	nap Score	NCBI gi	Position	Gene Id	Contig.Id	S E S
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1368 Bt1Gc1706 Bt1G3459	4717-3932	g2293262	971	979	1.40E-98	73	100	TRANSFERASE SUBUNIT ALPHA [Bacillus subtilis] (AF008220) acetyl-CoA carboxylase subunit [Bacillus
								subtilis]
1368 Bt1Gc1706 Bt1G3460	5577-3921	g1945709	142	183	3.10E-14	38	4	(Z94043) hypothetical protein [Bacillus subtilis]
1368 Bt1Gc1706 Bt1G3461	6881-5652	g2293261	1727	1589	3.10E-163	&	100	(AF008220) YtsJ [Bacillus subtilis]
1368 Bt1Gc1706 Bt1G3462	10314-6985	g2293260	2861	2951	1.50E-307	51	100	(AF008220) DNA-polymerase III alpha-chain [Bacillus subtilis]
1368 Bt1Gc1706 Bt1G3463	10767-11267	g2293326	296	243	1.40E-20	36	99	(AF008220) YtrI [Bacillus subtilis]
1368 Bt1Gc1706 Bt1G3464	12254-11316	g2293259	1029	1035	1.60E-104	61	100	(AF008220) YtqI [Bacillus subtilis]
1368 Bt1Gc1706 Bt1G3465	12414-12713	g2293325	203	205	1.40E-16	42	99	(AF008220) YtpI [Bacillus subtilis]
1368 Bt1Gc1706 Bt1G3466	14514-13201	g2293258	1567	1586	6.50E-163	68	100	(AF008220) YtoI [Bacillus subtilis]
1368 Bt1Gc1706 Bt1G3467	15643-14325	g2649315	468	538	7.40E-52	46	100	(AE001017) conserved hypothetical protein [Archaeoglobus fulgidus]
1368 Bt1Gc1706 Bt1G3468	15796-16884	g2633757	1135	1141	9.40E-116	58	100	(Z99111) similar to Xaa-Pro dipeptidase [Bacillus subtilis]
1369 Bt1Gc1715 Bt1G3469	392-63	g1172708	297	304	4.60E-27	55	99	PTS SYSTEM, CELLOBIOSE-SPECIFIC IIA
			7 m					COMPONENT (EIIA-CEL) (CELLOBIOSE-PERMEASE IIA COMPONENT) (PHOSPHOTRANSFERASE
				•				ENZYME II, A COMPONENT) (EIII-CEL) [Bacillus subtilis]
1369 Bt1Gc1715 Bt1G3470	1881-174	g1172710	933	978	1.80E-98	4	100	PTS SYSTEM, CELLOBIOSE-SPECIFIC IIC COMPONENT (EIIC-CEL) (CELLOBIOSE-PERMEASE
								IIC COMPONENT) (PHOSPHOTRANSFERASE ENZYME II, C COMPONENT) [Bacillus.subtilis]
1309 BUIGET/13 BUIG54/1	1061-007	82499992	310	310	2.50E-28	2	ý	COMPONENT (EIIB-CEL) (CELLOBIOSE-PERMEASE
								IIB COMPONENT) (PHOSPHOTRANSFERASE ENZYME II, B COMPONENT) [Bacillus
1369 Bt1Gc1715 Bt1G3472	4184-2838	g2499991	1515	1394	1.50E-142	65	100	Stearothermophilus J PTS SYSTEM, CELLOBIOSE-SPECIFIC IIC
		٠						COMPONENT (EIIC-CEL) (CELLOBIOSE-PERMEASE)
							·	ENZYME II, C COMPONENT) [Bacillus
1369 Bt1Gc1715 Bt1G3473	4492-4187	·g1172709	332	343	3.40E-31	62	99	PTS SYSTEM, CELLOBIOSE-SPECIFIC IIB
*****								COMPONENT (EIIB-CEL) (CELLOBIOSE-PERMEASE

[Bacillus subtilis] (D86947) chemotactic transducer [Pseudomonas	83	27	2.30E-50	524	427	2626835	15165-13077 g2626835	Bt1G3493	1370 Bt1Gc1717
3'REGION (ORF2) [Staphylococcus xylosus] METHYL-ACCEPTING CHEMOTAXIS PROTEIN MCPC	100 1	31	1.20E-74	753	756	g1708962	15068-13077	Bt1G3492	1370 Bt1Gc1717
(Z99120) yusU [Bacillus subtilis] HYPOTHETICAL 12.1 KD PROTEIN IN GLKA	99 (58 54	9.20E-22 8.70E-26	254 292	283 271	g2635789 g3023656	12681-12397 g2635789 12990-12676 g3023656	Bt1G3490 Bt1G3491	1370 Bt1Gc1717 1370 Bt1Gc1717
(D84432) DacF [Bacillus subtilis] (Z99120) similar to proline dehydrogenase [Bacillus subtilis]	74	34 60	7.60E-27 4.70E-98	302 974	267 974	g1303997 g2635781	10321-12466 g1303997 12073-11168 g2635781	Bt1G3488 Bt1G3489	1370 Bt1Gc1717 1370 Bt1Gc1717
(Z99120) similar to 3-hydroxyacyl-CoA dehydrogenase	100	83	1.50E-291	2800	2861	g2635780	9910-7455	c1717 Bt1G3487	1370 Bt1Gc1717
subtilis] (Z99120) similar to acetyl-CoA C-acyltransferase [Bacillus	100	79	1.20E-161	1574	1564	g2635779	7430-6261	c1717 Bt1G3486	1370 Bt1Gc1717
L-lactate dehydrogenase (EC 1.1.1.27) - Bacillus subtilis [] (AF027868) YobT [Bacillus subtilis] (Z99120) similar to butyryl-CoA dehydrogenase [Bacillus	100 100	46 59 76	6.70E-74 5.20E-67 1.20E-216	746 681 2093	785 739 2319	g80325 g2619046 g2635778	1332-382 3989-3291 5913-4132	Bt1Gc1717 Bt1G3483 Bt1Gc1717 Bt1G3484 Bt1Gc1717 Bt1G3485	1370 Bt1Gc1717 1370 Bt1Gc1717 1370 Bt1Gc1717
presumably containing the arsenate-arsenite-antimony transmembrane channel [Bacillus subtilis] THIAMIN BIOSYNTHESIS PROTEIN THIC [Bacillus subtilis]	100	78	2.70E-253	2439	2468	g3041750	18377-16617 g3041750	Bt1G3482	
(AF008220) YthA [Bacillus subtilis] (AF008220) YthB [Bacillus subtilis] (Z93767) similar to ArsB integral membrane protein	100	58 58	1.30E-68 3.20E-115	696	814 1337	g/293292 g/29338	13688-14722 g2293292 15178-16506 g1929338		
[Streptomyces coelicolor] (Z99106) ydaJ [Bacillus subtilis]	100	35	5.40E-49	511	455 1173	g2632727	12816-10764 g2632727	Bt1Gc1715 Bt1G3478 Bt1Gc1715 Bt1G3479	1369 Bt1C
INTERGENIC REGION (ORFB) [Bacillus subtilis] (AL049763) putative transcriptional regulator	100	38	2.90E-50	523	502	g4757086	9835-10716	Bt1Gc1715 Bt1G3477	1369 Bt1C
[Bacillus stearothermophilus] (Z99107) similar to hypothetical proteins [Bacillus subtilis] HYPOTHETICAL 84.1 KD PROTEIN IN HEMY-GLTT	99	35 32	4.20E-17 1.60E-58	210 601	150 766	g2633029 g2851530	7279-7650 9691-7700	Bt1Gc1715 Bt1G3475 Bt1Gc1715 Bt1G3476	1369 Bt1C 1369 Bt1C
IIB COMPONENT) (PHOSPHOTRANSFERASE ENZYME II, B COMPONENT) [Bacillus subtilis] PROTON/SODIUM-GLUTAMATE SYMPORT PROTEIN (GLUTAMATE-ASPARTATE CARRIER PROTEIN)	100	25	4.70E-34	370	328	g121467	4711-5952 g121467	1369 Bt1Gc1715 Bt1G3474	1369 Bt10
NCBI gi description	%	% % Ident Cvrg	Table 1 BlastP- Prob	BlastP Score	aat Bi	NCBI gi	Position	Contig Id Gene Id	SEQ ID Co

	د								
_	52 100	9.90E-73		735	g2293167	14638-13862	18 Bt1G3518	1373 Bt1Gc1718	_
_	58 100	1.60E-111	101	1153	g2293168	13794-12695	18 Bt1G3517	1373 Bt1Gc1718	_
_	26 100	4.80E-33	256	300	g1498192	10365-11693' g1498192	18 Bt1G3516	1373 Bt1Gc1718	
0 (D28119) putative [Pseudomonas aeruginosa]	26 100	4.80E-33	256	300	g1498192	10365-11693	18 Bt1G3515	1373 Bt1Gc1718	_
	53 100	4.80E-64	653	624	ģ1945684	9547-10209	18 Bt1G3514	1373 Bt1Gc1718	_
	57 99	1.30E-56	583	564	g2293301	8618-9202	18 Bt1G3513	1373 Bt1Gc1718	
_	77 100	2.50E-138	. 1354	1344	g2293302	7662-8627			
0 HYPOTHETICAL 39.9 KD PROTEIN IN AMYLASE	57 100	5.80E-84	841	993	g418336	6985-5951	18 Bt1G3511	1373 Bt1Gc1718	_
	49 100	3.70E-66	673	643	g2293177	5601-4843	18 Bt1G3510	1373 Bt1Gc1718	_
9 (Y14080) hypothetical protein [Bacillus subtilis]	62 99	2.90E-50	523	511	g2226150	2022-1531	18 Bt1G3509	1373 Bt1Gc1718	
0 (Y14080) hypothetical protein [Bacillus subtilis]	77 100	2.10E-118		1577	g2226151	1530-307			_
(nuoL1) [Rickettsia prowazekii]									
_	37 53	1.60E-39	425	399	g3861319	11328-15150	16 Bt1G3507	1372 Bt1Gc1716	_
5 [Synechococcus sp.]					. •				
NADH	40 100	8.50E-112	927	1061	g400640	10436-12289	16 Bt1G3506	1372 Bt1Gc1716	
	37 .74	6.10E-96	954	863	g2633696	5032-1783	16 Bt1G3505	1372 Bt1Gc1716	_
-	53 100	9.90E-73	735	693	.g730100	16849-17640	20 Bt1G3504	1371 Bt1Gc1720	_
subtilis]					. <i>i</i>				
	72 100	0	3768	3878	g2635168	16554-13618	20 Bt1G3503	1371 Bt1Gc1720	· <u>~</u>
INTE									
	45 99	9.40E-29	320	354	g3915537	13602-13123	20 Bt1G3502	1371 Bt1Gc1720	_
	28 100	1.70E-22	261	201	g2226218	11884-11189	20 Bt1G3501	1371 Bt1Gc1720	
O ALANINE DEHYDROGENASE [Bacillus sphaericus]	73 100	3.70E-144	1409	1400	g118515	10723-9611	20 Bt1G3500	1371 Bt1Gc1720	_
	52 100	1.90E-101	1006	1328	g2116759	9489-8110		1371 Bt1Gc1720	
[Synechocystis sp.]									•
[Bacillus subtilis] 9 TRANSCRIPTIONAL REPRESSOR SMTB HOMOLOG	36 99	1.30E-22	262	247	g2495428	7199-6804	20 Bt1G3498	1371 Bt1Gc1720	:
(Z991	66 100	3.30E-239	2306	2322	g2635862	6772-4415	20 Bt1G3497	1371 Bt1Gc1720	_
	70 100	6.70E-193	1869	1842	g2635670	2739-4208	20 Bt1G3496	1371 Bt1Gc1720	<u></u>
	97 100	1.20E-211	2046	2128	g473955	2546-1275	20 Bt1G3495	1371 Bt1Gc1720	_
aeruginosa] 8 (L77763) neutral protease A [Bacillus thuringiensis]	72 68	2.00E-121	834	1101	g2239276	842-1	20 Bt1G3494	1371 Bt1Gc1720	
				Score	•			·	
NCBI gi description	nt Cvrg	BlastP- % Prob Ident	BlastP Score		NCBI gi	Position	ld : Gene Id	SEQ ID Contig Id	<i>'</i>
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1374 Bt1Gc1719	4 Bt1Gc1719	4 Bt1Gc1719	4 Bt1Gc1719	4 Bt1Gc1719 4 Bt1Gc1719	4 Bt1Gc1719	4 Bt1Gc1719	4 Bt1Gc1719	4 Bt1Gc1719		1374 Bt1Gc1719	4 Bt1Gc1719	4 Bt1Gc1719	4 Bt1Gc1719	3 Bt1Gc1718	3 Bt1Gc1718	3 Bt1Gc1718		3 Bt1Gc1718	Contigues
	Bt1G3536	Bt1G3535	Bt1G3534	Bt1G3532 Bt1G3533	Bt1G3531	Bt1G3530	Bt1G3529	Bt1G3528		Bt1G3527	Bt1G3526	Bt1G3525	Bt1G3524	Bt1G3523	Bt1G3522	Bt1G3521		Bt1G3519	
Bt1G3537 12043-13341 g2506579	10962-11897	9957-10763	8931-9938	7337-7558 6884-8922	7055-6363	6234-5398	4366-5334	4128-3370		3187-1881	1722-1294	1395-847	443-779	20714-19636	18447-17938	17938-17335	15222-17057	14713-15225	
g2506579	g2633168	g1706796	g1706797	g2636297 g1706795	g732380	g732378	g730415	g732376		g461913	g1175629	g732366	g732365		g2293241			g2293300	q
1659	205	922	954	321 963	574	845	1278	932		879	470	259	322	1525	551	305	568	648	Score
1666	332	871	693	321 957	402	861	1278	941		974	380	242	340	1501	555	337	657	576	Score
2.20E-171	5.00E-30	3.80E-87	2.80E-68	7.30E-29 3.00E-96	1.90E-60	4.40E-86	2.80E-130	1.50E-94		4.70E-98	4.10E-35	1.70E-20	7.10E-31	6.70E-154	1.20E-53	1.50E-30	1.80E-64	7.00E-56	Prob
71	28	65	54	77 51	54	61	79	67		46	61	40	· 2	84	61	35	35	7 5 .	Ident
100	100	100	100	99 100	100.	100	100	100		100	99	99	99	90	99.	99	100	99	Cyrg
HYPOTHETICAL 51.0 KD PROTEIN IN PTA 3'REGION	(Z99108) similar to iron(III) dicitrate transport permease		_		[Bacillus subtilis] HYPOTHETICAL 29.7 KD PROTEIN IN PTA 3'REGION [Bacillus subtilis]	PROTEIN 43) (VEG43) [Bacillus subtilis] HYPOTHETICAL 31.4 KD PROTEIN IN PTA 3'REGION	PROBABLE PHOSPHATE ACETYLTRANSFERASE (PHOSPHOTRANSACETYL ASE) (VEGETATIVE	HYPOTHETICAL 29.5 KD PROTEIN IN ROCC-PTA	CARBOXYPEPTIDASE) (CPASE) (PBP5) [Bacillus	D-ALANYL-D-ALANINE CARBOXYPEPTIDASE	INTERGENIC REGION [Bacillus subtilis] CELL WALL HYDROLASE CWLJ [Bacillus subtilis]	HYPOTHETICAL 20.3 KD PROTEIN IN UNG-ROCA	HYPOTHETICAL 12.0 KD PROTEIN IN UNG-ROCA	protein [Bacillus subtilis] (AF008220) SAM synthase [Bacillus subtilis]	(AF008220) putative molibdenum cofactor biosynthesis	(Z99111) molybdopterin-guanine dinucleotide biosynthesis	(D63999) hypothetical protein [Synechocystis sp.]	(AF008220) YtoA [Bacillus subtilis]	0

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Contig Id		1374 Bt1Gc1719		Bt1Gc1719	1374 Bt1Gc1719	Bt1Gc1719	1374 BHG-1710	0:100111	Bt1Gc1719	Bt1Gc1719	Bt1Gc1721		Bt1Gc1/21	Bt1Gc1721	Bt1Gc1722		Bt1Gc1722	Bt1Gc1722	Bt1Gc1722	1376 Bt1Gc1722		Bt1Gc1722	1376 Bt1Gc1722	
Gene Id		Bt1G3538		Bt1G3539	Bt1G3540	Bt1G3541	B+1(354)		Bt1G3543	Bt1G3544	Bt1G3545		Bt1G3546	Bt1G3548	Bt1G3549	-	Bt1G3550	Bt1G3551	Bt1G3552	Bt1G3553		Bt1G3554	Bt1G3555	
Position	·	13572-13387 g1565237		13685-14347	14382-14897	16978-15047	17700_17080 @1731048	11100	17952-18356	18365-19456 g4914624	3447-4426		9870-117482	11990-13839	156-1067		2475-3557	3566-4204	4223-5413	5432-5893		7401-6399	10108-8939	
NCBI gi		7 g1565237				7 g1565240	0 01731048	61701000	5 g2116754	5 g4914624	g4530143		g4126620			•	g132565	g132633	g121062	g2497762	• ,.	g1705464	g115009	
aat_ nap Score	}	238		620	690	1740	434	ş	260	1071	383	· •	224	200	830		1005	656	1532	588		1012	776	
BlastP Score	· }	243		. 552	654	1741	453	į	283	920	354)	346	230	715		1006	618	1541	416		1031	789	
BlastP- Prob		1.40E-20		2.40E-53	3.80E-64	2.50E-179	7 50E_43		7.80E-25	2.50E-92	2.30E-32		8.80E-17	3.20E-19	1.30E-70		1.90E-101	2.50E-60	3.80E-158	6.30E-39		4.20E-104	1.90E-78	
% % Ident Cvrg		79		53	76	53	2		39	57	41		ر ا ا					61	73	73		59	43	
Cvrg	}	98	•	100	99	100	3	į	99	92	100	<u>.</u>	57 00 100	79	. 100		100	100	100	99	٠.	100	100	
NCBI gi description	[Bacillus subtilis]	(Z80360) Unknown, highly similar to Pseudomonas putida	4-oxalocrotonate tautomerase [Bacillus subtilis]	(Z80360) Unknown [Bacillus subtilis]	(Z80360) Unknown [Bacillus subtilis]	(Z80360) unknown, highly similar to penicillin binding	protein [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis]	(D86418) YfmP [Bacillus subtilis]	(AJ009627) multidrug resistance transporter [Listeria	monocytogenes] (AF085222) putative scaffolding protein [Streptococcus	thermophilus bacteriophage DT1]	(AB016282) ORF34 [bacteriophage phi-105]	(AB016282) ORF37 [hacterionhage nhi-105]	HYPOTHETICAL 34.6 KD PROTEIN IN GLNQ-ANSR	INTERGENIC REGION [Bacillus subtilis]	RIBOFLAVIN-SPECIFIC DEAMINASE [Bacillus subtilis]	RIBOFLAVIN SYNTHASE ALPHA CHAIN [Bacillus subtilis]	BUTANONE 4-PHOSPHATE SYNTHASE (DHBP	6,7-DIMETHYL-8-RIBITYLLUMAZINE SYNTHASE	(RIBOFLAVIN SYNTHASE BETA CHAIN) [Bacillus	BIOTIN SYNTHASE (BIOTIN SYNTHETASE) [Bacillus subtilie]	8-AMINO-7-OXONONANOATE SYNTHASE (7-KETO-8-AMINO-PELARGONIC ACID SYNTHETASE) (7-KAP	SYNTHETASE) (L-ALANINEPIMELYL COA LIGASE) [Bacillus sphaericus]

Table 1

(AE000933) dTDP-glucose 4,6-dehydratase	63	4	8.00E-71	717	669	g2622920	647-1	Bt1G3571	1377 Bt1Gc1723	13
HYPOTHETICAL 34.0 KD PROTEIN IN GLNQ-ANSR	. 99	60	1.20E-99	989	964	g1731073	31495-30585	Bt1G3570	1376 Bt1Gc1722	. 13
	100	59	1.30E-63	649	671	g2337814	29700-29041 g2337814	Bt1G3569	1376 Bt/IGc1722	13
HOMOLOG 2 [Bacillus subtilis] (Y13937) putative YhaP protein [Bacillus subtilis]	100	72	2.00E-90	902	1072	g2337815	29022-28112	Bt1G3568	1376 Bt1Gc1722	13
	100	48	7.50E-59	604	4 2	g1709791	Bt1G3567 27263-28099 g1709791		1376 Bt1Gc1722	13
INTERGENIC REGION [Bacillus subtilis]										
	100	47	1.70E-75	761	706	g1731077	Bt1G3566 27097-25719 g1731077		1376 Bt1Gc1722	13
INTERGENIC REGION [Bacillus subtilis]						,				
	. 100	56	2.20E-68	694	730	g1731078	Bt1G3565 26125-25337 g1731078		1376 Bt1Gc1722	13
1.2.1.38) - Bacillus subtilis [Bacillus subtilis]			•			•				-
	100	53	4.90E-87	870	967	g80340	Bt1G3564 24893-23859 g80340		1376 Bt1Gc1722	ដ
SYNTHASE) (AGS) [Bacillus stearothermophilus]					•					
ACETYLTRANSFERASE (N-ACETYLGLUTAMATE										
, TRANSACETYLASE) (OATASE) / AMINO-ACID										
ACETYLTRANSFERASE) (ORNITHINE										
	100	60	2.10E-127	1251	1233	g584771	23854-22634 g584771	Bt1G3563	1376 Bt1Gc1722	13
PHOSPHOTRANSFERASE) [Bacillus stearothermophilus]										
(N-ACETYL-L-GLUTAMATE 5-					·					
ACETYLGLUTAMATE KINASE (NAG KINASE) (AGK)	100	48	4.70E-50	521	543	g584767	22619-21852	Bt1G3562	1376 Bt1Gc1722	
[Bacillus subtilis]						•				
ACETYLORNITHINE AMINOTRANSFERASE (ACOAT)	100	54	1.60E-97	969	1032	g543846	21852-20699 g543846	Bt1G3561	1376 Bt1Gc1722	13
ANABOLIC (OTCASE) [Bacillus subtilis]										
ORNITHINE CARBAMOYLTRANSFERASE,	100	58	7.70E-89	887	924	g129265	20682-19729	Bt1G3560	1376 Bt1Gc1722	13
(D83967) YfkN [Bacillus subtilis]	4	48	1.30E-173	1663	1450	g2626826	17072-15133 g2626826	.Bt1G3559	1376 Bt1Gc1722 Bt1G3559	13
NA(+)/H(+) ANTIPORTER [Bacillus firmus]	100	41	4.00E-92	918	1001	g3287970	13315-14694 g3287970	Bt1G3558	1376 Bt1Gc1722	13
AMINOTRANSFERASE) [Bacillus subtilis]				•						
AMINOTRANSFERASE) (DAPA										
DIAMINO-PELARGONIC ACID						,				
OXONONANOATE AMINOTRANSFERASE (7,8-										
ADENOSYLMETHIONINE-8-AMINO-7-	100	59	5.20E-138	1351	1416	g1705460	12163-10814	Bt1G3557	1376 Bt1Gc1722 Bt1G3557 12163-10814 g1705460	13
subtilis]						-				
	100	4	6.00E-50	520	495	g1705466	10814-10119 g1705466	Bt1G3556	1376 Bt1Gc1722	13
	,				Score					
NCBI gi description	Cvrg %	% % Ident Cvrg	BlastP- Prob	BlastP Score	nap	NCBI gi	Position	Gene Id	Q Contig Id	
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DIHYDROLIPOAMIDE SUCCINYLTRANSFERASE COMPONENT OF 2-OXOGLUTARATE	100	. 66	1.00E-125	1235	1347	g129041	7219-8475	Bt1G3593	1378 Bt1Gc1724
oxoglutarate dehydrogenase (lipoamide) (EC 1.2.4.2) - Bacillus subtilis []	100	53	1.10E-268	2584	2494	g1071810	4215-7040	Bt1G3592	1378 Bt1Gc1724
nypothetical protein MJ02/2 - Methanococcus Jannaschii [Methanococcus Jannaschii]	9	9/	5.30E-19	228	199	10787178	3443-3190	Bilussyl	13/8 BHGC1/24
(Y09946) transposase [Bacillus thuringiensis]	<u>4</u>	ន្	1.70E-27		280	g1749770	1-369	Bt1G3590	
MECA [Bacillus subtilis]									
NEGATIVE REGULATOR OF GENETIC COMPETENCE	70	51	3.90E-30	333	299		23784-23307	Bt1G3589	
(D88209) Pz-peptidase [Bacillus licheniformis]	<u>1</u> 00	70	4.60E-226	.2182	2312		23082-18475	Bt1G3588	
(Z99110) yjbH [Bacillus subtilis]	1 00	43	2.90E-50	523	591		17126-17956	Bt1G3587	1377 Bt1Gc1723
(Z99110) yjbI [Bacillus subtilis]	99	67	2.20E-43	458	438		16653-17051	Bt1G3586	1377 Bt1Gc1723
(Z99110) yjbK [Bacillus subtilis]	99	50	3.40E-40	. 428	429	g2633512	15907-16479	Bt1G3585	1377 Bt1Gc1723
(Z99110) yjbL [Bacillus subtilis]	99	39	1.80E-11	157	210	g2633513	15762-15394	Bt1G3584	1377 Bt1Gc1723
(Z99110) similar to GTP pyrophosphokinase [Bacillus	100	65	3.00E-71	721	707	g2633514	15363-14728 g2633514	Bt1G3583	1377 Bt1Gc1723 Bt1G3583
(Z99110) similar to hypothetical proteins [Bacillus subtilis]	100	72	4.30E-95	946	1046	g2633515	14703-13906 g2633515	Bt1G3582	1377 Bt1Gc1723
INTERGENIC REGION [Bacillus subtilis]									
HYPOTHETICAL 31.5 KD PROTEIN IN MECA-TENA	100	54	5.40E-81	813	803		13843-12995	Bt1G3581	1377 Bt1Gc1723
(AB016282) ORF41 [bacteriophage phi-105]	31	30	2.10E-25	299	100	g4126627	12074-13384	Bt1G3580	1377 Bt1Gc1723
subtilis]						• •			
aeruginosaj (Z99110) similar to diadenosine tetraphosphatase [Bacillus	100	57	1.50E-76	771	753	g2633517	10697-11431 g2633517	Bt1G3579	1377 Bt1Gc1723
(AF14/448) rod-shape-determining protein [Pseudomonas	00	32	6.40E-46	482	337	g488/204	9451-10617	B11G35/8	13// Bt1Gc1/23
(D90901) hypothetical protein [Synechocystis sp.]		40	3.60E-18	230	166	g1651989	8298-8649	Bt1G3577	
[Mycobacterium tuberculosis]	:	;) ! ! .					
[Micromonospora griseorubida] HYPOTHETICAL 23.1 KD PROTEIN CY277.20C	10	47	2.00E-51	534	528	g2829568	7667-8281	Bt1G3576	1377 Bt1Gc1723
(D16097) mycinamicinIII O-metyltransferase	100	46	1.70E-61	629	574	g303643	6500-7345	Bt1G3575	1377 Bt1Gc1723
(AF071085) Orfde14 [Enterococcus faecalis]	50	. 30	4.60E-19	232	159	g3608402	4015-1843	Bt1G3574	
PROTEIN SPSI [Bacillus subtilis]						;			•
[Enterococcus faecalis] SPORE COAT POLYSACCHARIDE BIOSYNTHESIS	100	52	4.30E-63	644	668	g730818	1951-1220	Bt1G3573	1377 Bt1Gc1723
(AF071085) dTDP-4-dehydrorhamnose 3,5-epimerase	99	56	2.20E-52	543	517	g3608395	1205-648	Bt1G3572	1377 Bt1Gc1723
[Methanobacterium thermoautotrophicum]						,			-
NCBI gi description	% Cvrg	% Ident (BlastP- Prob l	BlastP Score	nap Score	NCBI gi	Position	Gene Id	SEQ ID Contig Id NO

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1380	1380 1380	i379 1379	1379 1379	1379 1379 1379	13/9	1379	1379	1379	1379 1379	1379 1379	1378 1378 1379	NO SEQ
Bt1Gc1728	Bt1Gc1728 Bt1Gc1728	Bt1Gc1726 Bt1Gc1726		Bt1Gc1726 Bt1Gc1726 Bt1Gc1726					1379 Bt1Gc1726 1379 Bt1Gc1726	Bt1Gc1726 Bt1Gc1726	Bt1Gc1724 Bt1Gc1724 Bt1Gc1726	Contig Id
Bt1G3616	Bt1G3614 Bt1G3615	Bt1G3612 Bt1G3613	Bt1G3610 Bt1G3611	Bt1G3607 Bt1G3608 Bt1G3609	Bt1G3606	Bt1G3604 Bt1G3605	Bt1G3602	Bt1G3601	Bt1G3599 Bt1G3600	Bt1G3597 Bt1G3598	Bt1G3594 Bt1G3595 Bt1G3596	Gene Id
6156-5331	4134-454 5357-4530	14675-15325 14669-16048	13497-11889 13615-14325	11345-10731 11657-12964	9885-9379	8001-9146	5708-6562	4951-3885	2823-3053 3510-3161	1088-684 2545-1199	11311-12462 15151-14771 1-1338	Position
g1709791	g1041115 g2635853	g4584090 g2500998	g4584088 g4584089	g585341 g1934782 g3258354	g2982937	g1934789 g585644	g2125/91 g1945659	g1934791	g2635634 g3123231	g1934804 g120717	g4218544 g2497400 g1934805	NCBI gi
398	892 1093	1157 401	711 1184	379 636 973	317	760 913	901	1073	284 274	316 2051	. 171 187 242	aat_ nap Score
428	993 1095	1157 507	754 1184	391 636 878	345	778 929	866	1088	223 270	339 2006	377 229 311	BlastP Score
3.40E-40	4.50E-100 7.00E-111	1.90E-117 1.50E-90	9.60E-75 2.60E-120	2.80E-36 3.10E-62 6.90E-88	2.10E-31	2.70E-77 2.70E-93	9.30E-45 1.30E-86	3.90E-110	1.80E-18 1.90E-23	9.10E-31 2.00E-207	6.10E-34 4.10E-19 8.40E-28	BlastP- Prob
33	39 74	100 32		59 57,	37	45 3	6 8	8	74 52	85	39	% % Ident Cvrg
100	100	100	81 100	100	99	100	100	100	99	99 100	50 57 43	Cvrg
metabolite dehydrogenase [Bacillus subtigis] PYRROLINE-5-CARBOXYLATE REDUCTASE HOMOLOG 2 [Bacillus subtilis]	(D78016) TRAC [Enterococcus faecalis] (Z99121) alternate gene name: yvsB; similar to plant-	cereus] (AJ010128) glycyl-tRNA synthetase [Bacillus cereus] PROBABLE GLYCYL-TRNA SYNTHETASE (GLYCINE-TRNA LIGASE) (GLYRS) [Methanococcus jannaschii]	horikoshii] (AJ010128) hypothetical protein [Bacillus cereus] (AJ010128) DNA alkylation repair enzyme [Bacillus	KINASE-ASSOCIATED PROTEIN B [Bacillus subtilis] (Z93933) unknown [Bacillus subtilis] (AP000007) 424aa long hypothetical protein [Pyrococcus	(AE000679) superoxide dismutase (Cw/Zn) [Aquitex aeolicus]	(Z93934) unknown [Bacillus subtilis] PUTATIVE AMINOTRANSFERASE B [Bacillus subtilis]	(Z93934) unknown [Bacillus subtilis] (Z94043) hypothetical protein [Bacillus subtilis]		(Z99120) yuzA [Bacillus subtilis] GENERAL STRESS PROTEIN 13 (GSP13) [Bacillus	(Z93936) unknown [Bacillus subtilis] GLUCOSE-6-PHOSPHATE ISOMERASE A (GPI A) (PHOSPHOGLUCOSE ISOMERASE A) [Bacillus stearothermonbilus]	(AJ011500) gra-orf12 [Streptomyces violaceoruber] HYPOTHETICAL TRANSPOSASE-LIKE PROTEIN HI1721 [Haemophilus influenzae Rd] (Z93936) unknown [Bacillus subtilis]	NCBI gi description

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SEQ ID Contig Id Gene Id	Position	NCBI gi	aat_	BlastP Score	BlastP- Prob	% Ident	Cvrg	NCBI gi description
1380 Bt1Gc1728 Bt1G3617	7302-6448	g1805461	935	820	9.70E-82	62	100	(D50453) homologues to hypothetical 30.5 kDa protein gdhl
						٠.		5' region of B. megateruim [Bacillus subtilis]
1380 Bt1Gc1728 Bt1G3618	8175-7613	g2635852	538	506	1.80E-48	59	. 99	(Z99121) alternate gene name: yvsC; similar to molybdenum transport permasse [Pacillus subtilis]
1380 Bt1Gc1728 Bt1G3619	9070-8289	g2635851	700	737	6.10E-73	57	100	(Z99121) alternate gene name: yvsD; similar to molybdate-
))	}	binding protein [Bacillus subtilis]
Bt1Gc1728		g1772644	1133	1056	9.50E-107	74	100	(U62055) orfR gene product [Bacillus subtilis]
1380 Bt1Gc1728 Bt1G3621	11586-12554	g1934659	1177	1088	3.90E-110	68	100	(U93876) hypothetical protein YrdR [Bacillus subtilis]
1380 Bt1Gc1728 Bt1G3622	13210-13806	g2226210	515	518	9.80E-50	52	99	(Y14082) hypothetical protein [Bacillus subtilis]
1380 Bt1Gc1728 Bt1G3623	14282-15256	g2633758	701	743	1.40E-73	4	100	(Z99111) similar to transcriptional regulator (LacI family)
1380 Bt1Gc1728 Bt1G3624 15871-16707		g1783243	766	722	2.40E-71	50	100	(D83026) homologous to jojC gene product (B. subtilis;
			ļ					prf:2111327a); hypothetical [Bacillus subtilis]
Bt1Gc1/28 Bt1G3625	1/83/-18685	g1789981	796	806	3.00E-80	52	100	
1381 B(10C1/2/ B(103020	1-1000	81001240	202	0,40	J.JUE-J0	9/	9	PRODUCT IN MANY BACTERIA. [Bacillus subtilis]
1381 Bt1Gc1727 Bt1G3627	6530-1675	g4835822	1317	1216	1.10E-123	54	100	(AF102174) glycine betaine transporter BetL [Listeria
1381 BriGc1727 BriG3628	10487-13255	62636153	2310	2406	8 40F-250	ś	<u> </u>	monocytogenes] (790172) similar to SNE2 beliege [Bacillus subtilis]
Bt1Gc1727		g2633808	1012	776	4.50E-77	52	100	(Z99111) similar to hypothetical proteins [Bacillus subtilis]
1381 Bt1Gc1727 Bt1G3630	19161-18475	g2832800	718	729	4.30E-72	. 62	100	(AJ223978) putative ABC transporter, YvrO [Bacillus
			į) ;		, · ·		subtilis]
1381 BtlGc1/2/ BtlG3631	20321-19164	g2633806	471	343	3.40E-31	29	100	(299111) similar to hypothetical proteins from B. subtilis [Bacillus subtilis]
1381 Bt1Gc1727 Bt1G3632	21925-20600	g585209	1011	929	2.70E-93	46	100	GLYCEROL-3-PHOSPHATE TRANSPORTER (G-3-P
D.10-1706	137 133	2000	3	3		5	•	TRANSPORTER) (G-3-P PERMEASE) [Bacillus subtilis]
1382 Bri Gc1725 Bri G3634	3073-2420	6730508	1000	1000	1 00E-02	73	3 5	1-DABBOI INE 2-CABBOAAL VIE DEHADBOGENVOE
- ;		-	,	,		;		(PSC DEHYDROGENASE) [Bacillus subtilis]
1382 Bt1Gc1725 Bt1G3635	8217-6211	g2632976	2359	2287	3.40E-237	69	100	(Z99107) similar to DNA ligase [Bacillus subtilis]
1382 Bt1Gc1725 Bt1G3636	10485-8236	g2577965	2592	2590	2.70E-269	68	100	(Y15254) PcrA protein [Bacillus subtilis]
1382 Bt1Gc1725 Bt1G3637	11184-10498	g2577964	793	811	8.70E-81	63	100	(Y15254) PcrB protein [Bacillus subtilis]
1382 Bt1Gc1725 Bt1G3638	14452-13190	g131611	1376	1399	4.30E-143	63	100	PHOSPHORIBOSYLAMINEGLYCINE LIGASE (GARS)
y t								(GLYCINAMIDE RIBONUCLEOTIDE SYNTHETASE) (PHOSPHORIBOSYLGLYCINAMIDE SYNTHETASE)

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1384 Bt1Gc1730	1384 Bt1Gc1730	1384 Bt1Gc1730 1384 Bt1Gc1730	1384 Bt1Gc1730 1384 Bt1Gc1730	1384 Bt1Gc1730			1384 Bt1Gc1730	1383 Bt1Gc1730		1383 Bt1Gc1729	1383 Bt1Gc1729	1383 Bt1Gc1729	·1383 Bt1Gc1729	1383 Bt1Gc1729	1383 Bt1Gc1729	1383 Bt1Gc1729	SEQ ID Contig Id NO
	1730	1730 1730	1730 1730				1730	1730	3	1729	729	729	729		729		3 Id
Bt1G3671	Bt1G3670	Bt1G3668 Bt1G3669	Bt1G3666 Bt1G3667	Bt1G3665			Bt1G3664	Bt1G3663		Bt1G3661	Bt1G3660	Bt1G3659	Bt1G3658	Bt1G3657	Bt1G3656	Bt1G3655	Gene Id
8691-7480	7455-6949	6446-5820 6853-6536	3393-3990 5660-4164	3252-1504	35		1420-545	1-484	1011	14022-15447	13129-15364	12850-15085	12850-14803	12850-14524	12850-14245	12850-13963 g135254	Position
£1934829	g2129134	g2635704 g2635705	g3169326 g2635702	g2851553			g585225	g2635700		g732329	g135254	g135254	g135254	g135254	g135254	g135254	NCBI gi
1256	116	533 310	341 1481	1938			1229	376	3	525	86	221	224	221	219	220	nap Score
1230	229	505 196	389 1472	1958		•	1161	252	8	691	130	260	269	277	269	270	BlastP Score
3.50E-125	4.10E-19	2.30E-48 1.30E-15	4.60E-36 7.90E-151	2.50E-202			7.10E-118	1.40E-80 1.50E-21		4.50E-68	9.00E-08	3.80E-22	3.30E-23	3.60E-24	3.30E-23	2.60E-23	BlastP- Prob
5 82	30	\$ 51 5 54	5 38 1 58	96			8 1	47		35	3 54	. 69	55	96	1 77	67	% Ident
100	99	100	. 100	100	•		100	81	<u>;</u>	100	=	19	26	21	. 16	. 20	%Cvrg
[Methanococcus jannaschii]) (Z93939) unknown [Bacillus subtilis]				PROBABLE PHOSPHOMANNOMUTASE (PMM) [Bacillus subtilis]	URIDYLYLTRANSFERASE) (URIDINE DIPHOSPHOGLUCOSE PYROPHOSPHORYLASE) (GENERAL STRESS PROTEIN 33) (GSP33) [Bacillus	URIDYLYLTRANSFERASE (UDP-GLUCOSE PYROPHOSPHORYLASE) (UDPGP) (ALPHA-D-GLUCOSYL-1-PHOSPHATE	UTPC	(AE000433) IS130 putative transposase [Escherichia coli] (Z99120) similar to biotin metabolism [Bacillus subtilis]		ENZYME RES [Bacillus cereus] HYPOTHETICAL 49.5 KD PROTEIN IN DAE-TYRZ) TYPE III RESTRICTION-MODIFICATION SYSTEM ENZYME RES [Bacillus cereus]	NCBI gi description

Table 1

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N. V. C. L. M. M. D. M.	70		-	607	6113673	3003-1266	B+1G3605		1387
(A E010A06) hymothetical protein [Phodobacter canculatus]	22 75	3 SOE-20	2445 1.0	100	8178703	7648-329	B+1G3604	7 B+1Gc1733	1387
		-		3307	~107.497.5	22/1/2 20/201	D+1C2C02		120
(Y11478) probably site-specific recombinase of the	35 100	2.80E-59	608 2	521	g1865708	19441-21005	Bt1G3692	Bt1Gc1732	1386
(L44593) repressor protein [Lactococcus lactis phage BK5-	34 . 51	1.10E-16	206 1	120	g928836	17140-18847	Bt1G3691	Bt1Gc1732	1386
(D90903) hypothetical protein [Synechocystis sp.]				122	g1652146	11248-10047	Bt1G3690		1386
INTERGENIC REGION (ORF1) [Bacillus subtilis]								•	,
HYPOTHETICAL 24.0 KD PROTEIN IN PONA-COTD	31 100	4.50E-29	323 4	223	g732244	9373-8583	Bt1G3689	Bt1Gc1732	1386
(AB016282) ORF17 [bacteriophage phi-105]	39 99	7.30E-29	321 7	317	g4126654	7031-7444	Bt1G3688	6 Bt1Gc1732	1386
coelicolor]									•.
(AL023861) putative IS element transposase [Streptomyces	26 100	2.80E-20 :	240 2	169	g3218350	6002-5310	Bt1G3687	Bt1Gc1732	1386
(AF010496) hypothetical protein [Rhodobacter capsulatus]	28 100	3.00E-55	570 3	410	g3128374	2338-4026	Bt1G3686	6 Bt1Gc1732	1386
(AF011378) unknown [Bacteriophage sk1]	31 99	7.00E-08	123 7	79	g2392827	1157-1437	Bt1G3685	Bt1Gc1732	1386
(AF047044) putative transposase [Anabaena PCC7120]	20 72	7.90E-17	208 7	117	g3005554	1-685	Bt1G3684	Bt1Gc1732	1386
[Haemophilus influenzae Rd]									
PUTATIVE GLYCOSYL TRANSFERASE HI0868	36 100	9.20E-38	405 9	387	g3123077	23836-23087	Bt1G3683	Bt1Gc1731	1385
INTERGENIC REGION [Bacillus subtilis]									
HYPOTHETICAL 42.0 KD PROTEIN IN DAPB-PAPS	25 90	5.90E-27	303 5	171	g1730929	24094-22005	Bt1G3682	Bt1Gc1731	1385
thermoautotrophicum				:	•				
(AE000878) putative membrane protein [Methanobacterium	23 72	2.00E-60	649 2	321	g2622173	14392-19516	Bt1G3681	Bt1Gc1731	1385
[Cryptosporidium parvum]									
(AF068065) GP900; mucin-like glycoprotein	24 31		199 3	124	g4063042	4975-24258	Bt1G3680	Bt1Gc1731	1385
(Y09476) YitH [Bacillus subtilis]	34 100	3.40E-40	428 3	409	g2145400	5089-4235	Bt1G3679	Bt1Gc1731	1385
INTERGENIC REGION [Bacillus subtilis]	٠.				(
HYPOTHETICAL 14.7 KD PROTEIN IN BCSA-DEGR	49 99	1.90E-30 ′	336 1	330	g1730902	1876-2271	Bt1G3678	Bt1Gc1731	1385
lactis				i					-
hypothetical protein, 35K - Lactococcus lactis [Lactococcus	34 100	8.80E-49	509 8	436	e282295	937-1809	Bt1G3677	Bt1Gc1731	1385
					,			٠.	
(X99978) ORF7; hydophobic protein [Lactobacillus	36 99	7.80E-25	283 7	251	g1922884	761-267	Bt1G3676	Bt1Gc1731	1385
(AB002150) YbbK [Bacillus subtilis]	52 99	8.00E-39	415 8	410	g1256140	23563-24015	Bt1G3675	Bt1Gc1730	1384
(AF027868) DNA-binding protein YobU [Bacillus subtilis]	37 99	4.50E-29	323 4	271	g2619047	20695-21171	Bt1G3674	Bt1Gc1730	1384
NF-180 - sea lamprey [Petromyzon marinus]	26 64	5.40E-20 2	256 5	199	g2133786	26525-7739	Bt1G3673	Bt1Gc1730	1384
(Z93939) thioredoxine reductase [Bacillus subtilis]	74 100	3.20E-131	1287 3.2	1319	g1934830	9147-10142	Bt1G3672	Bt1Gc1730	1384
				Score				•	NO
NCBI gl description	$\overline{}$	Prob Ident			NCBI gi	Position	Gene Id	Contig Id	Ħ,
	, °	RiggtP_ %	Riser PRI	aat					SEQ

(Z99111) molybdopterin biosynthesis protein [Bacillus subtilis]	63 100	3.30E-136	1334	1392	g2633799	7647-6358	Bt1G3717	1389 Bt1Gc1734 Bt1G3717	138
(ORF 2) [Bacillus megaterium] (Z99111) molybdopterin converting factor (subunit 2)	61 99	2.60E-49	514	505	g2633801	5870-5405	Bt1G3716	9 Bt1Gc1734	1389
HYPOTHETICAL 30.5 KD PROTEIN IN GDHI 5'REGION	82 100	2.10E-111	1100	1227	g731355	4626-3772	Bt1G3715	9 Bt1Gc1734	1389
OF MYXOCOCCUS XANTHUS. [Bacillus subtilis] GLUCOSE 1-DEHYDROGENASE [Bacillus megaterium]	88 100	8.50E-122	1198	1198	g729328	3755-2973	Bt1G3714	9 Bt1Gc1734	1389
(AB001488) SIMILAR TO TRANSCRIPTION FACTOR	55 99	1.00E-31	348	412	g1881322	2418-2876	Bt1G3713	9 Bt1Gc1734	1389
(Z99113) yndB [Bacillus subtilis]	63 99	1.80E-48	506	495	g2634157	958-524	Bt1G3712		1389
(D90903) hypothetical protein [Synechocystis sp.]	22 84	4.90E-23	266	116	g1652146	26292-23643			1388
INTERGENIC REGION (ORF1) [Bacillus subtilis]									
HYPOTHETICAL 24.0 KD PROTEIN IN PONA-COTD	32 100	1.00E-29	329	227		23214-22413	Bt1G3710	8 Bt1Gc1739	1388
(AB016282) ORF17 [bacteriophage phi-105]	39 99	4.50E-29	323	319	g4126654	20871-21284	Bt1G3709	8 Bt1Gc1739	1388
thermophilus bacteriophage DT1]									
(AF085222) putative scaffolding protein [Streptococcus	31 94	4.40E-22	257	251	g4530143	16049-14381	Bt1G3708	8 Bt1Gc1739	1388
(AF009630) 116 [bacteriophage bIL 170]	25 100	1.40E-61	440	307	g3282276	10634-7189	Bt1G3707	8 Bt1Gc1739	1388
(X97918) gene 18.1 [Bacteriophage SPP1]	30 47	5.60E-38	411	342	g2764873	8733-6448	Bt1G3706	8 Bt1Gc1739	1388
bacteriophage O1205]							•		
(U88974) ORF42 [Streptococcus thermophilus temperate	42 6	4.40E-08	138	62	g2444121	4302-2520	Bt1G3705	8 Bt1Gc1739	1388
(AF047044) putative transposase [Anabaena PCC7120]	24 36	1.50E-08	135	97	g3005554	1-353	Bt1G3704	8 Bt1Gc1739	1388
thermophilus bacteriophage DT1]									
(AF085222) putative scaffolding protein [Streptococcus	27 88	7.50E-20	236	166	g4530143	20831-19158	Bt1G3703	7 Bt1Gc1733	1387
phage BK5-T (fragment) []									
hypothetical protein 1 (bpi 5' region) - Lactococcus lactis	42 99	6.40E-14	180	126	g320251	20104-19580 g320251	Bt1G3702	7 Bt1Gc1733	1387
Promove aminowe prowar approximations program to the program of th			t	į					į
promoter inhihitor protein BpI - I actoroccius lactis phage		2 20E-29	9CF	258	6320252	19546-18938	Bt1G3701		1387
(AF097906) myosin heavy chain [Rana catesbeiana]	•	5.10E-13	193	75	g4249699	16154-13023	Bt1G3700		1387
(AJ006589) gp43 [Bacteriophage phi-C31]	27 57	1.40E-32	36 4	240	g3947462	16615-12628	Bt1G3699	7 Bt1Gc1733	1387
(AB016282) ORF37 [bacteriophage phi-105]	31 79	2.50E-19	231	194	g4126623	12383-10534	Bt1G3698	7 Bt1Gc1733	138
(D13377) XpaF1 protein [Bacillus licheniformis]	34 95	6.10E-09	133	97	g2160199	3770-3473	Bt1G3697	7 Bt1Gc1733	1387
3'REGION (ORFD) [Clostridium perfringens]									
HYPOTHETICAL 14.9 KD PROTEIN IN NAGH	32 99	7.10E-15	189	180	g141088	3428-3036	Bt1G3696	7 Bt1Gc1733	1387
			•	•					
CWI:A PRECURSOR (CELL WALL HYDROLASE)					•				
NCBI gi description	% % Ident Cvrg	Prob Idei	Score	₹D.	NCBI gi	Position	Gene Id	Contig Id	8 E
			714D	aat	`		وغرد	<i>i</i>	SEQ

	cereus] (AJ010139) polysaccharide deacetylase-like protein	100	. 99	9.10E-134	1311	1375	g4584141	Bt1G3737 22682-21903 g4584141	Bt1G3737	1390 Bt1Gc1736	1390
	subtilis] (AJ010139) DNA alkylation repair enzyme [Bacillus	100	93	1.70E-146	1431	1431	g4584142	21840-20980 g4584142	Bt1G3736	Bt1Gc1736	1390
	(PSEUDOURIDYLATE SYNTHASE I) (PSEUDOURIDINE SYNTHASE I) (URACIL HYDROLYASE) [Bacillus sp.] (Z99108) similar to RNA methyltransferase [Bacillus	100	66	1.00E-157	1537	1632	g2633126	Bt1G3735 20972-19581		Bt1Gc1736	1390
•	(D/8193) positive regulatory protein [Bacillus subtilis] TRNA PSEUDOURIDINE SYNTHASE A	100	62	9.30E-77	773	813	g1064808 g3915176	19530-18796	Bt1G3734	Bt1Gc1736	1390
	AMINO-ACID PERMEASE ROCE [Bacillus subtilis]	100	58	3.40E-143	1400	1510	g730601	16355-14952			1390
•	INTERGENIC REGION [Bacillus subtilis] (Z99112) similar to acetylornithine deacetylase [Bacillus	100	63	6.00E-153	1492	1485	g2633908	14829-13552	Bt1G3731	Bt1Gc1736	1390
	HYPOTHETICAL 34.0 KD PROTEIN IN NPRE-PYCA	100	56	3.50E-93	928	928	g2494800	10319-11245	Bt1G3730	Bt1Gc1736	1390
	(D86417) YfIF [Bacillus subtilis]		61	1.60E-120	1186	1358	g2443228	9993-8491			1390
• •	biosynthesis protein [Pyrococcus horikoshii] (Z99112) penicillin-binding protein [Bacillus subtilis]	100	48	1.20E-85	857	82 5	g2634067	8237-6999	Bt1G3728	Bt1Gc1736	1390
	(AP000002) 455aa long hypothetical Vi polysaccharide	100	41	5.80E-84	841	776	g3256832	4722-2534	Bt1G3727	Bt1Gc1736	1390
-	pyruvate formate-lyase (pfl) homolog - Haemophilus influenzae (strain Rd KW20) Haemophilus influenzae Rd1	26	55	9.20E-50	521	578	g1075196	597-1	Bt1G3726	Bt1Gc1736	1390
	LEUCYL-TRNA SYNTHETASE (LEUCINETRNA LIGASE) (LEURS) [Bacillus subtilis]	71	77	9.50E-242	2330	2411	g3123286	25860-24146	Bt1G3725	Bt1Gc1734	1389
• ••	(AF067645) spore germination protein GerIB [Bacillus cereus]	100	92	2.60E-168	1637	1738	g3290176	19518-20609	Bt1G3724	Bt1Gc1734	1389
_	(AF067645) spore germination protein GerIA [Bacillus cereus]	100	95	1.00E-237	2273	3186	g3290175	17276-19498	Bt1G3723	Bt1Gc1734	1389
	(AF067645) homoserine O-acetyltransferase homolog	100	99	1.70E-123	1214	1214	g3290174	17124-16435	Bt1G3722	Bt1Gc1734	Į389
	(Z99119) similar to hypothetical proteins from B. subtilis	99	66	7.20E-22	255	243	g2635515	15462-15259	Bt1G3721	Bt1Gc1734	1389
	(299110) similar to thiamin biosynthesis [Bacillus subtilis] NARA PROTEIN [Bacillus subtilis]	100	57 57	8.70E-97	962	1008	g/30103	10306-9287	Bt1G3720	Bt1Gc1734	1389
	(D88802) E. coli moaC protein; P30747 (368) [Bacillus subtilis]	•	68	9.20E-54	556	583	g1945112	7731-8248			1389
	NCBI gi description	Cvrg	% Ident	BlastP- Prob	BlastP Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	NO DEC
in the		_	,	TADICT		•					}. }

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SEQ ID Contig Id Gene Id NO	Position	NCBI gi	aat_ nap Score	BlastP Score	BlastP- Prob	% % Ident Cvrg	%	NCBI gi description
1390 Bt1Gc1736 Bt1G3738	Bt1G3738 24330-22813 g	g417009	2132	2155	3.30E-223	77	100	[Bacillus cereus] FUMARATE HYDRATASE CLASS I, AEROBIC
								(FUMARASE) [Bacillus stearothermophilus]
1390 Bt1Gc1736 Bt1G3739	24929-25390 g	g2626813	680	680	6.70E-67	80	99	(D83967) YfkA [Bacillus subtilis]
1390 Bt1Gc1736 Bt1G3740	25589-26047 g	g2626814	576	578	4.30E-56	71	99	(D83967) YfkB [Bacillus subtilis]
1390 Bt1Gc1736 Bt1G3741	26892-26095 g	g2626816	721	725	1.10E-71	54	100	(D83967) YfkD [Bacillus subtilis]
1390 Bt1Gc1736 Bt1G3742	28021-26972 g	g2626817	1306	1010	7.10E-102	73	100	(D83967) YfkE [Bacillus subtilis]
1390 Bt1Gc1736 Bt1G3743	28194-28616 g	g2829682	285	294	5.30E-26	3 8	99	GENERAL STRESS PROTEIN 26 (GSP26) [Bacillus
•								subtilis]
1390 Bt1Gc1736 Bt1G3744	30162-29677 g	g2632514	340	262	1.30E-22	41	99	(Z99105) similar to alkaline phosphatase [Bacillus subtilis]
1391 Bt1Gc1738 Bt1G3745	6139-5237 g	g1749770	735	735	9.90E-73	51	100	(Y09946) transposase [Bacillus thuringiensis]
1391 Bt1Gc1738 Bt1G3746	6570-7908 g	g4103625	1050	927	4.50E-93	49	100	(AF026470) gluconate permease [Pseudomonas aeruginosa]
1391 Bt1Gc1738 Bt1G3747	11833-10575 g	g709994	598	586	6.10E-57	39	.100	(D14399) hypothetical protein [Bacillus subtilis]
1391 Bt1Gc1738 Bt1G3748	12582-11811 g	g1176995	835	851	5.00E-85	61	100	HYPOTHETICAL ABC TRANSPORTER ATP-BINDING
1391 Bt1Gc1738 Bt1G3749	16606-12618 g1731300	1731300	857	757	4.60E-75	36	<u> </u>	HYPOTHETICAL 70.5 KD PROTEIN IN IDH 3'REGION
٠,								[Bacillus subtilis]
1391 Bt1Gc1738 Bt1G3750	17723-16743	g1176994	566	631	1.00E-61	38	100	HYPOTHETICAL SENSOR-LIKE HISTIDINE KINASE
 								IN IDH 3'REGION [Bacillus subtilis]
1391 Bt1Gc1738 Bt1G3751	18421-17735 g	g1176993	657	661	6.90E-65	55	100	HYPOTHETICAL 26.6 KD SENSORY TRANSDUCTION
								PROTEIN IN IDH 3'REGION [Bacillus subtilis]
1391 Bt1Gc1738 Bt1G3752	18631-18900 g	g2635892	333	336	1.90E-30	67	99	(Z99121) similar to transcriptional regulator (ArsR family)
1391 Bt1Gc1738 Bt1G3753	21117-20248 6	01881325	1311	1211	3 60F-173	77	3	(AROO1488) PROBABI E HTH ARAC FAMILY OF
		·						
1391 Bt1Gc1738 Bt1G3754	21427-22185 g	g2293177	707	712	2.70E-70	54	100	(AF008220) transporter [Bacillus subtilis]
1391 Bt1Gc1738 Bt1G3755	29515-28628 ε	g728972	582	602	1.20E-58	39	100	BMRU PROTEIN [Bacillus subtilis]
1391 Bt1Gc1738 Bt1G3756	30230-29691 g	g585264	447	460	1.40E-43	46	99	HYPOXANTHINE-GUANINE
	,	•						PHOSPHORIBOSYLTRANSFERASE (HGPRT)
1391 Bt1Gc1738 Bt1G3757	32538-33239 g	g1575577	400	390	3.60E-36	39	100	(HGPRTASE) [Bacillus subtilis] (U67196) DNA-binding response regulator [Thermotoga
	-	-			•			maritima]
	32736-35074 8	g466195	306	372	7.30E-34	35	41	SENSOR PROTEIN RESE [Bacillus subtilis]
1392 Bt1Gc1735 Bt1G3759	3545-497 g	g4062401	1252	1414	1.10E-144	47	93	
•				•			٠	protein in bcr 5'region. [Escherichia coli]

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1397 I 1398 I	1396 E	1395 E	1394 E	1393 E	1393 E		1393 E	1393 E	1393 E	1393 E	1393 E	1393 E	1392 E	1392 E	1392 E	1 7461		1392 E		SEO
Bt1Gc1763 Bt1Gc1767	Bt1Gc1765	Bt1Gc1762	Bt1Gc1758	Bt1Gc1737 Bt1Gc1737	Bt1Gc1737	Bt1Gc1737	Bt1Gc1737	Bt1Gc1737	Bt1Gc1737	Bt1Gc1737	Bt1Gc1737	Bt1Gc1737	Bt1Gc1735	Bt1Gc1735	Bt1Gc1735	667 130119	Bt1Gc1735	Bt1Gc1735	Contig Id	
1763 1767	1765		1758	1737 1737	1737	1737	1737		1737	1737		1737	1735	1735	1735			1735	g Id	•
Bt1G3780 Bt1G3781	Bt1G3779	Bt1G3778	Bt1G3777	Bt1G3775 Bt1G3776	Bt1G3774	Bt1G3772	Bt1G3771	Bt1G3770	Bt1G3769	Bt1G3768	Bt1G3767	Bt1G3766	Bt1G3765	Bt1G3764	Bt1G3763	20/07/19	Bt1G3761	Bt1G3760	Gene Id	
3780 3781	3779	3778	3777		3774	3772	3771		3769	3768	3767	3766	3765	3764	3763	20/02	3761	3760	e Id	
270 47	.	2	.	20607-19509 21742-20755	1845	1677	1522	13770-12046	9943	781:	364	16	11562	8840	8073	723.	371	336	Po	
270-117 47-478	1-222	212-1	1-282	7-195(2-207:	18459-17776	16771-15335	15228-14038)-120	9943-11574	7813-8709	3644-1671	1656-1	11562-10234	8846-9980	8072-8539	/235-3632	3714-3348	3362-2421	Position	
g5	6 2	6 2	<u>81</u>	35 g2;								8 39								
g595780 g1934652	g3123300	g3290175	g1724002	g2331287 g3122850	g2618833 g2618833	g2618834	g2618836	g1176951	g2769708	g732377	g3123297	g3915204	g731354	g2618997	g2635598	##10C1B	g3757815	g129184	NCBI gi	-
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122 478	191	244	139	1560 1195	752	1325	654	2189	955	642	2821	2286	1290	531	242	C##7	285	538	· • • • • • • • • • • • • • • • • • • •	99t 1
141 397	208	265	159	1491 1,169	757	1349	624	2199	852	649	2499	2287	1306	545	270	4747	337	462	Score	
8.7 6.5	7.1	5.7	2.9	7.60	4.6	8.50	5.7	7.20	3.9	1.3	1.20	3.40	3.10	1.3	1.9	1.00	1.5	. <u>4</u>	Prob	
8.70E-10 6.50E-37	7.10E-17	5.70E-22	2.90E-11	7.60E-153 1.00E-118	4.00E-69 4.60E-75	8.50E-138	5.70E-61	7.20E-228	3.90E-85	1.30E-63	1.20E-259	3.40E-237	3.10E-133	1.30E-52	1.90E-23	1.00E-201	1.50E-30	8.40E-44	÷ 5	
60	56	78	4	81 68	y 2	53	36	72	37	45	83	79	55	33	34		. 46	39	% Ident	?
) 27	5 23	=	30	100	100		100	100	100	100	100	58	100	100	99	I		100	0	₹ .
							7	_					•	_						
PROTEIN 35) (GSP35) [Bacillus subtilis] (U13871) lacZ alpha peptide [Cloning vector] (U93876) amino acid transporter [Bacillus subtilis]	cereus] THIOREDOXIN REDUCTASE (GENERAL STRESS	[Bacillus subtilis] (AF067645) spore germination protein GerIA	[Bacillus firmus] HYPOTHETICAL ABC TRANSPORTER ATP-BIND PROTEIN 2 IN GLPD-CSPB INTERGENIC REGION	Subunis] (AF013188) release factor 2 [Bacillus subtilis] PREPROTEIN TRANSLOCASE SECA SUBUNIT	(AF01/113) cell division protein [Bacillus subtilis] (AF017113) cell division ATP-binding protein [Ba	(AF017113) putative protease [Bacillus subtilis]	(AF017113) YyjD [Bacillus subtilis]	[Staphylococcus aureus] HYPOTHETICAL ABC	(U82085) pristinamycin resistance protein VgaB	[Bacillus subtilis] HYPOTHETICAI IN MMR-PTA IN	EXCINUCLEASE ABC SUBUNIT B (DINA PROTEIN)	INTERGENIC REGION (ORF8) [Bacillus subtilis] EXCINUCLEASE ABC SUBUNIT A [Bacillus subtilis]	НҮРОП	F0278	9119	IRANSPOSASE FOR INSERTION SEQUENCE ELEMENT IS231C [Bacillus thuringiensis]	(AF042861) putative OppB [Treponema denticola]	OLIGOPEPTIDE TRANSPORT SYSTEM PERMEASE PROTEIN OPPC [Bacillus subtilis]		
N 35)) lac2) ami	DOX	subti 45) sj	HETION N 2 II	188) release factor 2 [Bacillus subtilis] OTEIN TRANSLOCASE SECA SUBU	113) cell division protein [Bacillus subtilis] 113) cell division ATP-binding protein [Bacillus	13) p	13) Y	HETIC) prist	ıs subtilis] [HETICAL TRANSCRIPTIONAL REGULATOR R-PTA INTERGENIC REGION [Bacillus subtilis	JCLE,	CLE.	THETICAL 50.8 KD PROTEIN IN SRFA4-SFP	68) tr) simi	NT IS	61) p	PEPTIDE TRANSPORT SY IN OPPC [Bacillus subtilis]		
(GSF alpha no aci	N P	lis] pore g	NGLI CAL '	lease TRA	ell div	utativ ;:	ָלָ קלי ביי	CAL L	inam		ASE /	ASE /	CAL:	anscri	lar to	231C	utativ	PC (H	_	•
a pept d trar	EDUC	çermi	ABC	facto	ision/	e prot	Bacil	ABC Sens	ycin r	R R A	ABC (ABC S	50.8 K	ption	cyste	Back	e Opp	RANS Bacille	NCBI gi description	
Bacilide [TAS	nation	TRAI	r 2 [I OCA:	ATP.	ease	llus su	TRA:	esista	SCR SCR	SUBL	SUBL SUBL	D PF	antit	ine di	illus	B [1	SPOR us sut	gi de	
llus su Cloni er [B	E (GI	prote	NSPO NTE	3acill SE SE	in Le bindi	Baci	ibtilis	NSPO	nce p	REGI	TIN		OTE	ermin	oxyge		repor	TSY Silis]	scrip	
ng ve acillu	NER	in G	RGE	us sut	ing pr	llus s)	RTE	rotein		B (DI	Sacillı A [B	Z	ator	nase	piens	lema	STEN	tion	
ctor]	ALS	erlA	R ATI	otilis] SUBU	otein	ubtili:	rs suo	RATI	Vga	, REC	NA F	is sub acillu	SRF	Bacil	Bac		dentic	A PEI		
tilis]	TRE	[Bacillus	P-BIN	NIT	tilis] [Bac	. <u> </u>	CILIS	P-BIN	Φ.	}ULA llus sı	ROT	tilis] s subt	A4-S	llus sı	illus s	į.	ola]	ZME/		
	SS	llus	IS firmus] ITHETICAL ABC TRANSPORTER ATP-BINDING IN 2 IN GLPD-CSPB INTERGENIC REGION		illus			lococcus aureus THETICAL ABC TRANSPORTER ATP-BINDING THE TOTAL STREET OF TRANSPORTER ATP-BINDING		[Bacillus subtilis] HYPOTHETICAL TRANSCRIPTIONAL REGULATOR IN MMR-PTA INTERGENIC REGION [Bacillus subtilis]	EIN)	ilis]	FP	(AF027868) transcription antiterminator [Bacillus subtilis]	(Z99119) similar to cysteine dioxygenase [Bacillus subtilis]			SE		
			, 4 7		•		- '	ч.						•	ş	-		٠.	•	

Table 1

	1421 Bt1Gc1991 Bt1G3808 1-381 g2	1420 Bt1Gc1989 Bt1G3807 117-323 g4	1419 Bt1Gc1979 Bt1G3806 1-436 g1	1418 Bt1Gc1983 Bt1G3805 515-1 g3	1417 Bt1Gc1981 Bt1G3804 1-224 g2	Bt1Gc1960 Bt1G3803 1-508		1415 Bt1Gc1917 Bt1G3802 169-1 g3	1414 Bt1Gc1894 Bt1G3801 61-299 g2		1413 Bt1Gc1898 Bt1G3800 1-235 g5	1412 Bt1Gc1896 Bt1G3799 274-62 g2		1411 Bt1Gc1891 Bt1G3798 534-1 g2	1410 Bt1Gc1876 Bt1G3796 397-1 g2	1409 Bt1Gc1866 Bt1G3795 267-1 g1		1408 Bt1Gc1869 Bt1G3794 41-488 g3		1407 Bt1Gc1862 Bt1G3792 1-388 g1	1406 Bt1Gc1849 Bt1G3791 281-1 g2	1405 Bt1Gc1845 Bt1G3790 416-1 g4			1404 Bt1Gc1771 Bt1G3789 1-276 g1	1403 Bt1Gc1832 Bt1G3788 137-1 g1				1400 Bt1Gc1822 Bt1G3784 13-231 g3	1399 Bt1Gc1804 Bt1G3783 122-1 g3	NO	Contig Id Gene Id Position	SEQ
	g2983101	g4584092	g118797	g3401988	g2117766	g2635490		g3290175	g2116759		g586902	g2984723	~	g2501426	g2619017	g143324		g3401958	,	g1172446	g2462090	g4323583			g1172949	g124464	g4584140	g4584140		g321919	g3128280		NCBI gi	
	128	172	152	242	65	462	. •	80	124		132	190	•	691	92	203		562		98	154	73			277	135	116	234		102	94	Score		aat t
	183	. 181	170	293	120	484		114	140		153	194		706	151	159	•	565		14	181	113			280	156	150	, 256		97	110		Score)!^e+D
	3.10E-13	5.00E-14	3.20E-12	2.10E-25	1.00E-06	3.90E-46		9.60E-06	9.10E-09		2.10E-10	2.10E-15		1.20E-69	8.00E-10	5.50E-11		1.00E-54		6.90E-09	3.60E-13	7.70E-06		٠	1.60E-24	3.20E-10	1.90E-10	5.70E-22		4.00E-05	1.00E-05		Prob	Dlac+D_
	32	54	33	3 8	42	59		46	36	ė	47	58		80	33	54		74		26	39	21			66	60	53	66		3 80 .	49		=	8
٠	22	99	38	34	- 8	53		••	17		22	99		41	23	23		82		18	19	30			55	7	21	29		50	1	(Cyrg	\$
aeolicus	(AE000690) acetolactate synthase large subunit [Aquifex	(AJ010129) hypothetical protein [Bacillus cereus]	DNA POLYMERASE III, BETA CHAIN [Bacillus subtilis].	Perfringolysin O []	orthinine aminotransferase rocD - Bacillus subtilis []	(Z99119) similar to hypothetical proteins [Bacillus subtilis]	cereus].	(AF067645) spore germination protein GerIA [Bacillus	(D86418) YfnA [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis]	cereus] HYPOTHETICAL 40.7 KD PROTEIN IN MECB-GLTX	(AF053927) probable spore germination protein F [Bacillus	[Bacillus stearothermophilus]	PYRIMIDINE-NUCLEOSIDE PHOSPHORYLASE	(AF027868) squalene-hopene cyclase [Bacillus subtilis]	(M37169) APase I [Bacillus licheniformis]	Repressor, Dimeric Form [Bacillus subtilis]	Pyrr, The Bacillus Subtilis Pyrimidine Biosynthetic Operon	(KETO-ACID FORMATE-LYASE) [Escherichia coli]	KETO-ACID FORMATE ACETYLTRANSFERASE	(Y11171) BC542A protein [Bacillus cereus]	(AF101076) maturase-related protein [Pseudomonas putida]	PROTEIN 300) (VEG300) [Bacillus subtilis]	ACCLIMATIZATION PROTEIN) (CAP) (VEGETATIVE	thuringiensis] 50S RIBOSOMAL PROTEIN L10 (BL5) (COLD	IMMUNE INHIBITOR A PRECURSOR [Bacillus	(AJ010139) fumarate hydratase [Bacillus cereus]	(AJ010139) fumarate hydratase [Bacillus cereus]	plasmid NTP16 []	protein [Rhodobacter capsulatus] hypothetical 16.9K protein - Salmonella typhimurium	(AF0104		NCBI gi description	

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Table 1

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1443 Bt1Gc2141 Bt1 1444 Bt1Gc2171 Bt1	1442 Bt1Gc2162 Bt1	1441 Bt1Gc2158 Bt1	1440 Bt1Gc2138 Bt1	1439 Bt1Gc2134 Bt1	1438 Bt1Gc2122 Bt1	٠	1437 Bt1Gc2120 Bt1	1436 Bt1Gc2110 Bt1	1435 Bt1Gc2117 Bt1	1434 Bt1Gc2111 Bt1	1433 Bt1Gc2102 Bt1		1432 Bt1Gc2081 Bt1		1431 Bt1Gc2077 Bt1	1430 Bt1Gc2067 Bt1		1429 Bt1Gc2060 Bt1	1428 Bt1Gc2057 Bt1	Bt1Gc2029	 	1426 Bt1Gc2038 Bt1			1425 Bt1Gc2028 Bt1	1424 Bt1Gc2027 Bt1	23 Bt1Gc2025	• •	1422 Bt1Gc1992 Bt1	NO	SEQ ID Contig Id G
Bt1G3831 Bt1G3832	Bt1G3830	Bt1G3829	Bt1G3828	Bt1G3827	Bt1G3826		Bt1G3825	Bt1G3824	Bt1G3823	Bt1G3822	Bt1G3820		Bt1G3819	٠	Bt1G3818	Bt1G3817		Bt1G3816	Bt1G3815	Bt1G3814		Bt1G3813			Bt1G3812	Bt1G3811	Bt1G3810		Bt1G3809		Gene Id
441-220 541-1	1-280	328-1	316-1	340-1	270-1		371-1	1-423	243-1	1-352	1-214		1-195		160-507	255-1		249-1	186-1	397-1	• •	1-212		į	423-117	1-491	1-167		309-1		Position
g1001739 g3122117	g473955	g2612901	g586880	g1934652	g118515	(g1730918	g1220104	g3914289	g2633515	g4699725		g1787795	1	g2632987	g1001657		g1881268	g118516	g1945649		g2827439		9	9140583	g134774	g144312		g1708465	•	NCBI gi
104 222	105	188	171	197	148		419	162	126	155	140.		99		212	181	•	136	79	168		122			204	181	114		220	TD.	aat_]
119 212	156	222	189	208	165		.433	179	159	178	163		126		262	184		132	109	191		143		. [320	207	123		250		BlastP
4.60E-07 3.10E-16	1.40E-10	6.20E-18	7.10E-15	3.30E-16	1.10E-11		9.90E-41	2.30E-13	1.40E-10	1.00E-13	3.20E-11		1.70E-07		4.10E-22	3.30E-14		7.80E-08	1.40E-05	4.40E-15		3.70E-09			3 70F-18	2.30E-16	7.00E-08		1.60E-20	1100	r
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(D64004) ABC transporter [Synechocystis sp.] CELL DIVISION PROTEIN FTSH HOMOLOG [Bacillus	(D29674) DNA-binding protein [Lactobacillus sp.]	(AF015825) NADH dehydrogenase-like protein [Bacillus	PUR OPERON REPRESSOR [Bacillus subtilis]	(U93876) amino acid transporter [Bacillus subtilis]	ALANINE DEHYDROGENASE [Bacillus sphaericus]	INTERGENIC REGION [Bacillus subtilis]	carrier [Clostridium perfringens] HYPOTHETICAL 15.9 KD PROTEIN IN ILVD-THYB	(D49784) sodium-coupled branched-chain amino acid	TOPOISOMERASE IV SUBUNIT B [Bacillus subtilis]	(Z99110) similar to hypothetical proteins [Bacillus subtilis]	Bacillus Licheniformis Alpha-Amylase []	[Escherichia coli]	(AE000249) putative LACI-type transcriptional regulator	methyltransferase [Bacillus subtilis]	(Z99107) alternate gene name: yerS; similar to RNA	(D64002) acyl- [Synechocystis sp.]	HOMOLOG. [Bacillus subtilis]	stearothermophilus] (AB001488) ATP-DEPENDENT RNA HELICASE DEAD	ALANINE DEHYDROGENASE [Bacillus	(Z94043) hypothetical protein [Bacillus subtilis]	viscosus]	(AF043609) aluminum resistance protein [Arthrobacter	DEHYDROGENASE) [Bacillus subtilis]	REDITCTASE (TIDE-N-ACETYI MI IRAMATE		STAGE V SPORULATION PROTEIN E [Bacillus subtilis]	(J01566) 13.8 kd ORF [Plasmid CoIE1]	(VEGETATIVE PROTEIN 110) (VEG110) [Bacillus subtilis]	DIHYDROXY-ACID DEHYDRATASE (DAD)		NCBI gi description

Table 1

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	firmus]	ORF IS231C [Bacillus thuringiensis]	(AB013378) YhaQ [Bacillus halodurans]	lA-di	(version 2) - Bacillus subtilis []	PUTATIVE NAD	A-DI	RANS	BETA SUBUNIT) [Bacillus subtilis]	B001	(D13095) undefin		(D86418) YfmR [Bacillus subtilis]	(Z99122) ywzB [Bacillus subtilis]	(Z99120) fumarate hydratase [Bacillus subtilis]	(AF0156	(Z75208) hypothetical protein [Bacillus subtilis]	(M38482) HBsu protein [Artificial gene]	(D90907) hypothetical protein [Synechocystis sp.]	(AF022796) MoeB [Staphylococcus carnosus]	1832	(U26464) beta-galactosidase alpha [Cloning vector pZC320]	(Z93940) asparagine synthetase [Bacillus subtilis]	99117	ORF IS2	32952	subtilis]	(Y14080) hypothetical protein [Bacillus subtilis]	GMP SY	
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Contig Id Gene Id Position NCBI gi ant- ap nap score BlastP Prob Heat Cvrg 8 Bt1Gc2402 Bt1G3857 1-341 g1770035 180 217 6.30E-17 36 20 9 Bt1Gc2406 Bt1G3858 120-1 g586617 166 169 9.40E-13 83 35 9 Bt1Gc2406 Bt1G3859 534-254 g2633967 298 327 1.70E-29 69 28 9 Bt1Gc2410 Bt1G3860 389-1 g3121784 364 358 5.30E-32 59 19 1 Bt1Gc2412 Bt1G3861 558-1 g2633162 161 156 1.40E-10 25 44	subtilis]								٠,
Contig Id Gene Id Position NCBI gi ant-stream score BlastP score BlastP- % % % % % % Score % % % % Score % % % % Score Prob Ident Cvrg 8 Bt1Gc2402 Bt1G3857 1-341 g1770035 180 217 6.30E-17 36 20 36 169 9.40E-13 83 35 9 Bt1Gc2406 Bt1G3858 120-1 g586617 166 169 9.40E-13 83 35 9 Bt1Gc2406 Bt1G3859 534-254 g2633967 298 327 1.70E-29 69 28 9 Bt1Gc2410 Bt1G3860 389-1 g3121784 364 358 5.30E-32 59 19			1.40E-10	156	161	g2633162	558-1	Bt1G3861	1471 Bt1Gc2412
Contig Id Gene Id Position NCBI gi nap Score Prob Ident Cvrg 8 Bt1Gc2402 Bt1G3857 1-341 g1770035 180 217 6.30E-17 36 20 9 Bt1Gc2406 Bt1G3858 120-1 g586617 166 169 9.40E-13 83 35 9 Bt1Gc2406 Bt1G3859 534-254 g2633967 298 327 1.70E-29 69 28 9 Bt1Gc2410 Bt1G3860 389-1 g3121784 364 358 5.30E-32 59 19	PHOSPHOHYDROLASE [Clostridium								
Contig Id Gene Id Position NCBI gi nap Score Prob Ident Cvrg 8 Bt1Gc2402 Bt1G3857 1-341 g1770035 180 217 6.30E-17 36 20 9 Bt1Gc2406 Bt1G3858 120-1 g586617 166 169 9.40E-13 83 35 9 Bt1Gc2406 Bt1G3859 534-254 g2633967 298 327 1.70E-29 69 28			5.30E-32	358	364	g3121784	389-1	Bt1G3860	1470 Bt1Gc2410
Contig Id Gene Id Position NCBI gi nap Score Prob Ident Cyrg Score Score Prob Ident Cyrg Bt1Gc2402 Bt1G3857 1-341 g1770035 180 217 6.30E-17 36 20 Bt1Gc2406 Bt1G3858 120-1 g586617 166 169 9.40E-13 83 35	[Bacill (Z991		1.70E-29	327	298	g2633967	534-254	Bt1G3859	
Contig Id Gene Id Position NCBI gi nap Score Prob Ident Cvrg Score Stil Gc2402 Bt 1 G3857 1-341 g1770035 180 217 6.30E-17 36 20 (Z752)			9.40E-13	169	166	g586617	120-1	Bt1G3858	
Contig Id Gene Id Position NCBI gi nap Score Prob Ident Cvrg Score	(Z752)		6.30E-17	217	180	g1770035	1-341	Bt1G3857	
		% % ent Cvrg	'	BlastP Score		NCBI gi	Position	Gene Id	

Table 1

																						٠					
1506 Bt1Gc2639 Bt1G3897	1505 Bt1Gc2632 Bt1G3896	1504 Bt1Gc2631 Bt1G3895	1503 Bt1Gc2629 Bt1G3894	1502 Bt1Gc2622 Bt1G3893	1501 Bt1Gc2617 Bt1G3892	1500 Bt1Gc2613 Bt1G3891		1499 Bt1Gc2608 Bt1G3890	1498 Bt1Gc2594 Bt1G3889	1497 Bt1Gc2593 Bt1G3888	1496 Bt1Gc2570 Bt1G3887			1495 Bt1Gc2578 Bt1G3886		1494 Bt1Gc2577 Bt1G3885		1493 Bt1Gc2567 Bt1G3883	1492 Bt1Gc2563 Bt1G3882		1491 Bt1Gc2560 Bt1G3881	1490 Bt1Gc2559 Bt1G3880	•	1489 Bt1Gc2536 Bt1G3879	1488 Bt1Gc2532 Bt1G3878		SEQ ID Contig Id Gene Id NO
1-403	1-322	383-1	1-245	144	1-317	390-1		1-387	1-355	1-234	1-289	•		1-267		1-298		388-1	455-1		457-1	1-351	•	1-221	20-387		Position
 g1072499	g133732	g1730999	g4584142	g2337795	g1205984	g2634117	(e2635780	g1709733	g3821797	g132246	.•	4	g1730943		g130130		g2293298	g2635675		g1730908	g2500058		g1724002	g321919		NCBI gi
251	431	247	147	270	180	137		164	79	88	105			131	-	162	٠.	173	428		210	259		129	152	· !	aat_ nap Score
.214	<u>44</u>	305	190	217	127	149		229	152	139	74	•		149	,	134		213	386		139	252		139	166		BlastP Score
3.50E-17	1.40E-41	3.20E-26	5.60E-15	1.40E-16	5.60E-07	4.20E-10		6.20E-18	5.80E-10	1.40E-08	0.085			1.20E-10		5.70E-08		2.00E-17	9.50E-36		8.20E-09	1.80E-20		4.90E-09	2.00E-12		BlastP- Prob
4	79	49	50	48	47	32		ω Θ	37	46	31			39	,	43		41	· 56		35	54		4	ე დ	ı I	% % Ident Cvrg
37	76	17	28	16	12	41		16	15	14	26			60		17		56	41	•	40	16		25	83		% Cvrg
phospho-N-acetylmuramoyl-pentapeptide-transferase (EC 2.7.8.13) - Bacillus subtilis [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis] 30S RIBOSOMAL PROTEIN S12 (BS12) []	HYPOTHETICAL 79.2 KD PROTEIN IN PHOH-DGKA	(AJ010139) DNA alkylation repair enzyme [Bacillus	(Y13937) putative PacL protein [Bacillus subtilis]	(U33536) DNA: polymerase I [Bacillus stearothermophilus]	(Z99113) tRNA isopentenylpyrophosphate transferase [Bacillus subtilis]	[Bacillus subtilis]	(Z99120) similar to 3-hydroxyacyl-CoA dehydrogenase	PHOSPHOENOLPYRUVATE CARBOXYKINASE (ATP)	(D10594) chitinase D precursor [Bacillus circulans]	RECF PROTEIN [Bacillus subtilis]	subtilis]	IN UVRX-ILVA INTERGENIC REGION [Bacillus	HYPOTHETICAL TRANSCRIPTIONAL REGULATOR	PROTEIN PHOR [Bacillus subtilis]	ALKALINE PHOSPHATASE SYNTHESIS SENSOR	subtilis]	(AF008220) putative transcription regulator [Bacillus	(Z99120) similar to hypothetical proteins [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis]	FORMATE-LYASE) [Clostridium pasteurianum] HYPOTHETICAL 43.6 KD PROTEIN IN CPSD-METB	FORMATE ACETYLTRANSFERASE (PYRUVATE	PROTEIN 2 IN GLPD-CSPB INTERGENIC REGION	HYPOTHETICAL ABC TRANSPORTER ATP-BINDING	hypothetical 16.9K protein - Salmonella typhimurium	stearothermophilus]	NCBI gi description

Table 1

SUBTINS DIMETHYLADENOSINE TRANSFERASE (S- ADENOSYLMETHIONINE-6-N', N'-ADENOSYL(RRNA) DIMETHYLTRANSFERASE) (16S RRNA	35 45	2.70E-13	176	126	g585375	397-1	1526 Bt1Gc2765 Bt1G3918
	35 11	1.20E-12	182	118	g3915673	1-410	1525 Bt1Gc2755 Bt1G3917
	62 4	6.60E-10	156	153	g585920	1-159	1524 Bt1Gc2747 Bt1G3916
	46 16	4.90E-09	141	121	g1934606	188-1	1523 Bt1Gc2748 Bt1G3915
EPIMERASE (UDP-GLCNAC-2-EPIMERASE) [Bacillus							
	48 12	7.00E-06	112	99	g732324	1-136	1522 Bt1Gc2734 Bt1G3914
	45 25	4.20E-18	226	206	g1770026	353-1	
(X98626) SNF2 [Bacillus cereus]	35 21	5.50E-09	143	.	g1769944	348-1	_
SYNTHETASE) [Synechocystis sp.] (Z99112) similar to hypothetical proteins [Bacillus subtilis]	55 14	0.0055	87	149	g2634068	220-440	1519 Bt1Gc2725 Bt1G3911
	41 20	1.00E-17	224	159	g2494761	1-339	1518 Bt1Gc2700 Bt1G3909
(AJ010111) cytochrome caa3 oxidase assembly factor	48 26	2.70E-13	177	155	g4584150	7-248	1517 Bt1Gc2679 Bt1G3908
	36 19	3.10E-07	125	97	g2337799	1-238	1516 Bt1Gc2696 Bt1G3907
	39 26	0.032	77	76	g4584149	19-265	1515 Bt1Gc2685 Bt1G3906
	54 18	2.70E-19	237	214	g225559	1-255	1514 Bt1Gc2678 Bt1G3905
	56 23	6.30E-10	145	112	g3183561	185-1	1513 Bt1Gc2681 Bt1G3904
	50 15	7.10E-09	141	121	g2635763	1-215	1512 Bt1Gc2677 Bt1G3903
(AF027868) putative L-amino acid oxidase precursor	40 30	1.10E-24	284	240	g2619006	424-1	1511 Bt1Gc2656 Bt1G3902
(D87979) YfnK [Bacillus subtilis]	62 17	3.50E-12	169	135	g2116973	1-180	1510 Bt1Gc2652 Bt1G3901
	37 23	2.80E-10	154	146	g267497	549-233	1509 Bt1Gc2648 Bt1G3900
	27 50	4.20E-10	4	132	g1170998	330-1	1508 Bt1Gc2647 Bt1G3899
(GLUTAMATE-ASPARTATE CARRIER PROTEIN)	12 21	3.80E-32	332	419	g121400	/3-413	1307 Bt1Gc2641 Bt1G3898
	Ç		Score	Score		3	
NCBI gi description			BlastP	nap	NCBI gi	Position	SEQ D Contig Id Gene Id

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pSport1] (U01849) ORF1 [Trypanosoma brucei]	40 91	0.28	49	125	g436923	770-58	1545 Bt1Gc2910 Bt1G3938	1545 1
plasmid N1P16 [] (U12390) beta-galactosidase alpha peptide [cloning vector	38 81	6.60E-12	161	120	g531829	1-372	Bt1Gc2913 Bt1G3937	1544 I
[Bacillus subtilis] hypothetical 16.9K protein - Salmonella typhimurium	34 75	5.30E-10	143	79	g321919	357-18	Bt1Gc2878 Bt1G3935	1543 I
respiratory nitrate reductase alpha chain - Bacillus subtilis	63 4	3.30E-05	112	133	g2117582	145-1	Bt1Gc2865 Bt1G3934	1542 I
PRODUCT IN SYNECHOCYSTIS. [Bacillus subtilis]	31 30	1.205-09	141	120	81001333	33/-1	1341 611062639 61103933	1741 1
AMMONIA CHAIN) [Bacillus caldolyticus]		1 205 00	1	3	~1001766	267 1		1641
(CARBAMOYL-PHOSPHATE SYNTHETASE								
PYRIMIDINE-SPECIFIC: LARGE CHAIN		0.500	1,0	į	61100772	0		
CABBAMOVI BHOSBHATE SVAITHASE	40 45	3 50E-10	158	130	g1168777	305-1	Rt1Gc2862 Rt1G3032	1540 1
INTERGENIC REGION (ORF13) [Bacillus subtilis]		7301 30			1760100	1		
HYPOTHETICAL 31.7 KD PROTEIN IN GLTP-CWLJ	35 52	3.00E-16	202	132	g3183506	433-1	Bt1Gc2850 Bt1G3930	1538 I
catalase [Helianthus annuus]	26 24	3.60E-05	107	77	g1093432	1-351	Bt1Gc2843 Bt1G3929	1537 I
5'REGION (ORFB) [Bacillus subtilis]								
HYPOTHETICAL 27.6 KD PROTEIN IN ACUC	30 68	6.60E-12	161	180	g732300	505-1	Bt1Gc2842 Bt1G3928	1536 I
AMMONIA LIGASE) [Bacillus cereus]								
GLUTAMINE SYNTHETASE (GLUTAMATE	49 16	1.30E-11	166	125	g121357	1-212	Bt1Gc2835 Bt1G3927	1535 I
aeolicus		;	į	. :		1		
(AE000675) riboflavin specific deaminase [Aquifex		1.70E-05	108	77	£2982867	1-315		
(AJ010131) yfkH [Bacillus cereus]		5.50E-09	138	120	g4584097	145-1		
(AL021246) clpX [Mycobacterium tuberculosis]	68 · 14	2.90E-15	199	172	g2791497	1-182	Bt1Gc2797 Bt1G3924	1532 I
INTERGENIC REGION [Escherichia coli]					. (
I'RNA LIGASE) (I'HRRS) [Bacillus subtilis] HYPOTHETICAL 23.1 KD PROTEIN IN DMSC-PFLA	37 70	3.40E-15	192	160	g140338	452-1	1531 Bt1Gc2791 Bt1G3923	· 1531 I
THREONYL-TRNA SYNTHETASE I (THREONINE,	50 19	6.00E-20	246	270	g135175	371-1	Bt1Gc2789 Bt1G3922	1530 I
(U89796) chitinase [Bacillus thuringiensis]		1.00E-16	216	198	g2149596	182-1		1529 I
(Y11193) transcription antiterminator [Bacillus subtilis]	53 42	1.50E-23	271	343	g2154723	354-1		1528 I
(Z99117) similar to protease [Bacillus subtilis]		8.00E-06	. 110	110	. g2635181	212-108		
DIMETHYLTRANSFERASE) [Bacillus subtilis]			•	•				
DIMETHYLASE) (HIGH LEVEL KASUGAMYCIN			٠					٠.,
C. Control of the second of th	Ident Cvrg	Prob Id	Score	Score	ď			NO E
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coelicolor] 8 (D87979) YfnJ [Bacillus subtilis]	39	1.10E-08	144	100	g21,16974	1-271	3156 Bt1G3977	1580 Bt1Gc3156
	28 26	7.80E-07	122	82	g4204103	364-1	3149 Bt1G3976	1579 Bt1Gc3149
SYNTHASE (DXP SYNTHASE) [Bacillus subtilis]	. 0	4.005-17	7	2	81/31032	1-1		
		3.50E-10	340	3 5	61731057	334-1		
		\$ 30E-10	143	140	94584701	1-248		
		0 0001	<u> </u>	× ;	97331787	269-103		
ORF IS231C [Bacillus thuringiensis]	43 17	1.80E-06	119	& &	e225559	328-114	3135 Bt1G3972	1575 Bt1Gc3135
(TRANSCRIPTASE BETA CHAIN) (RNA POLYMERASE RETA SURUNIT) [Stanbulococcus aureus]							•	1
	47 12	3.10E-26	309	290	g1350849	410-1	3134 Bt1G3971	1574 Bt1Gc3134
ELEMENT IS232 [Insertion sequence IS232]				•	.*			
	43 21	4.60E-13	179	153	g2497382	268-1	3132 Bt1G3970	1573 Bt1Gc3132
	46 86	4.50E-29	323	312	g2633128	1-387	3115 Bt1G3969	1572 Bt1Gc3115
cereus					•	٠		
	50 46	2.30E-25	288	238	g4584142	1-395	3116 Bt1G3968	1571 Bt1Gc3116
					· ,•			
	49 20	8.60E-11	156	145	g3122129	205-1	3098 Bt1G3967	1570 Bt1Gc3098
INTERGENIC REGION [Bacillus subtilis]				-				
	43 18	7.20E-10	149	115	g732334	219-1	3086 Bt1G3966	1569 Bt1Gc3086
aeolicus]								-
(AE000679) superoxide dismutase (Cu/Zn) [Aquifex		3.20E-12	164	119	g2982937	396-8	3101 Bt1G3965	1568 Bt1Gc3101
	42 39	2.20E-20	241	187	g126054	1-379	3100 Bt1G3964	1567 Bt1Gc3100
			•		٠.			
		7.10E-21	257	238	g68538	1-272		1566 Bt1Gc3096
	39 29	5.30E-19	234	203	g2226226	1-398	3088 Bt1G3962	1565 Bt1Gc3088
DEHYDROGENASE) [Clostridium acetobutylicum]			٠					
					Ç.:			
	36 26	1.10E-10	156	112	g1703066	1-300		1564 Bt1Gc3087
[Bacillus subtilis] (AF008220) SAM synthase [Bacillus subtilis]	48 16	1.60E-10	155	133	g2293164	191-1	3070 Bt1G3960	1563 Bt1Gc3070
_	57 55	3.00E-32	353	447	g1805397	1-486	3060 Bt1G3959	1562 Bt1Gc3060
	52 30	5.20E-28		285	g473955	379-1		
protein [Bacillus subtilis]			•				•	
(Z92952) product similar to Staphilococcus aureus CapB	37 35	1.00E-08	133	117	g1894742	1-251	3055 Bt1G3957	1560 Bt1Gc3055
	Ident Cyrg	FF00	Score	Score	٠,٠			_
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NO BEQ	1601 E		1602 I	1603 I		604 I	1605 E	1606 F		1607 I	1608 I	1609 I		1610 I	1611 I		1612 I	1613 F			614 I		1615 I	1616 I		1617 1	1618 I	16191	
Contig Id	Bt1Gc3313		Bt1Gc3326	Bt1Gc3333		Bt1Gc3335	Bt1Gc3353	Bt1Gc3359		Bt1Gc3360	Bt1Gc3373	Bt1Gc3378		Bt1Gc3382	Bt1Gc3396		Bt1Gc3412	Rt1Gc3427	,		1614 Bt1Gc3430		Bt1Gc3434	Bt1Gc3445		Bt1Gc3450	Bt1Gc3463	Rt1Gc3494 Rt1G4017	
J Id																													+44
Gene Id	Bt1G3998		Bt1G3999	Bt1G4000		Bt1G4001	Bt1G4002	Bt1G4003		Bt1G4004	Bt1G4005	Bt1G4006		Bt1G4007	Bt1G4009		Bt1G4010	Rt1G4011			Bt1G4012		Bt1G4013	Bt1G4014		Bt1G4015	Bt1G4016	₽10.40	
	98		99	8		2	ຊ	8		2		8		07	9		10	=	,		12		13	14		15	16	i	_
Position	1-184		227-1	380-1		1-244	359-1	501-7		290-1	416-147	1-294		127-1	1-223		1-379	346-3	;		1-339		368-1	442-1		321-42	129-458	2	1-167
Z	g629265	•	g241	g246		g222	g227	g391	,-	g148683	g118262	g391	1	g406	g249	(g134396	g129521	q		g229		g283	g586866		g121190	g17(33	211
NCBI gi	265		g2415403	g2462099		g2226174	g2274944	g3915568		683	262	g3915989	-	g4063787	g2497382		396	<u>531</u>			g2293322		g2833392	866		190	g1706797		9//935/X
nap Score	99		152	285	·	85	134	185		327	398	189		136	129		263	130	į		118		188	177		139	249	124	_
BlastP Score	103		181	303		127		204		267	398	247		69	139		132	145			114		177	203	•	184	189	175	
BlastP- Prob	2.70E-05		2.90E-13	5.90E-27		2.50E-07	2.70E-10	5.20E-16		3.90E-23	5.10E-37	8.70E-21		0.067	1.00E-08		1.60E-07	3 30E-10	i		5.50E-06		1.30E-12	1.40E-15		2.40E-14	1.70E-14		/ / T
% Ident	45		49	52		33	33	33		63	90	58		63	49		50	ມ			33		43	သ အ		. 43	40	3	4/
% Cvrg	26	٠	17	55	•	17	30	4	•	52	99	24		27	17		15	٥	,		25		22	31		ა ა	32) 1	رد
		(frag	_		cereus]	_			HO											SXS						GLUC			
. ,	hyde c	(fragment) [Mycoplasma capricolum]	01577)908)	<u>s</u>	(Y14081) hypothetical protein [Bacillus subtilis]	(AJ000346) NapC protein [Enterococcus hirae]	HYPOTHETICAL 39.4 KD OXIDOREDUCTASE IN	M-MR	(M55343) open reading frame [Frankia sp.]	DNA-BINDING PROTEIN II (HB) (HU) [Bacillus	stearothermophilus] HYPOTHETICAL 4	INTERGENIC REGION (ORF4) [Bacillus subtilis]	00855	[Pseudomonas aeruginosa phage phi CTX] TRANSPOSASE FOR INSERTION SEOUENCE	ELEMENT IS232 [Insertion sequence IS232]	PREPROTEIN TRANSLOCASE SECA SUBUNIT	[Bacillus subtilis]	AMIDOTRANSFERASE COMPONENT II (ADC	SYNTHASE)/ ANTHRANILATE SYNTHASE	(AF008220) branch-chain amino acid	ilis]	HIO	HYPOTHETICAL 53.2 KD PROTEIN IN A	INTERGENIC REGION [Bacillus subtilis]	JCOSI	RICH		
	dehydi) [My	5) ace	sulfat		hypot	6) Nap	ETIC,	GA II	open	DINC	mophi ETIC/	NICI	O) orf	onas a OSAS	T ISZ	EN		RANS	SE)/	0) bra		ETIC	ETIC	NICI	EINE	ROM	[Bacillus subtilis]	
Z	ogena	copla	tylom	e trans		hetica	C pro	ÅL 39	NTER	readir	PRO	lus] AL 45	REGIO	14; si	erugu E FOI	32 [Ir	TRAN	2 S	FER.	ANT.	nch-c		AL 61	AL 53	REGI	III	ETR	s subt	<u> </u>
NCBI gi description	ase - N	sma c	itine	sport.		l prot	tein	.4 K	GEN	ng fra	TEIN	.3 KI	SN (6	milar	nosa p	sertic	OTS	Š	ASE C	P.A.	hain a		5 K	:	S E	ED D	ANSF	ilis]	
ji desc	Мусо _ј	aprico	deace	ATP-		ein []	Ente	000	CRE	me [I	H) II) PRC)RF4)	S Ot)hage ERTI	n seq	CASE	되 . 오 독	ÖMI		mino	٠.	PRO) PRC	Bacill	ISIVI	ÖRT		7
cripti	plasm	olum]	tylase	bindir		Bacill	госос	DOR	GIO	rank	B) (F	TEN	Ba	ene o	ON S The C	uence	SEC	Z T T	ÖNE	E SY	acid		JEI	TEI	us sul	ON P	PERI	2	_
) n	a capı		Bac	ıg pro		us sut	cus hi	EDU(V [Ba	ia sp.]	3	Z	cillus	f P2:t	EOU S	1S23	A SU	CE C	NTI	HIN	transp		Z		otilis]	ROTI	MEAS		0000
	ricolu		illus	tein		otilis]	rae]	CTAS	cillus		Bacill	ŘKA	subtil	ail co	ENCE CH	<u>2</u>	BUN	2 1 1	(AD	ASE	orter		DEC	(PAC		EIN B	SE PR	3	
	aldehyde dehydrogenase - Mycoplasma capricolum (SGC3)		(AF015775) acetylornitine deacetylase [Bacillus subtilis]	(Y10908) sulfate transport ATP-binding protein [Bacillus				EZ	HOM-MRGA INTERGENIC REGION [Bacillus subtilis]		us	nermophilus] THETICAL 45.3 KD PROTEIN IN PRKA-CSPB	<u>is</u>	(AB008550) orf14; similar to S gene of P2:tail completion	- 12	. •	Ŧ	DARA-AMINORENZOATE CYNTHACE CHITTAMINE	()		(AF008220) branch-chain amino acid transporter [Bacillus	1	HYPOTHETICAL 61.5 KD PROTEIN IN ADEC-PDHA	HYPOTHETICAL 53.2 KD PROTEIN IN XPAC-ABRB		OSE INHIBITED DIVISION PROTEIN B [Bacillus	FERRICHROME TRANSPORT PERMEASE PROTEIN	· 201	SS C
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1638 I	1635 H 1636 H 1637 H	1629 E 1630 E 1631 E 1632 E 1633 E 1634 E	1625 E 1626 E 1627 E 1628 E	1622 E 1623 E 1624 E	1620 E
1638 Bt1Gc3743 Bt1G4038	Bt1Gc3713 Bt1Gc3721 Bt1Gc3719	Bt1Gc3603 Bt1Gc3597 Bt1Gc3652 Bt1Gc3686 Bt1Gc3690 Bt1Gc3709	Bt1Gc3553 Bt1Gc3555 Bt1Gc3566 Bt1Gc3574		D Contig Id NO 1620 Bt1Gc3517
Bt1G4038	Bt1G4035 Bt1G4036 Bt1G4037	Bt1G4027 Bt1G4028 Bt1G4029 Bt1G4031 Bt1G4032 Bt1G4034	Bt1G4023 Bt1G4024 Bt1G4025 Bt1G4026	Bt1G4020 Bt1G4021 Bt1G4022	Gene Id Bt1G4018 Bt1G4019
530-89	380-145 1-310 1-278	245-1 256-1 212-464 66-405 258-1 1-301	1-411 337-1 1-333 1-210	1-469 219-1 372-1	Position 1-484 1-345
g1731017	g2829479 g1934609 g1169919	g130904 g1346795 g132246 g3688823 g2632519 g232186	g113346 g2127795 g1708794 g586885	g466137 g994736 g1169253	NCBI gi g2619056 g2633471
208	276 93 162	148 98 137 187 190	132 123 312 180	74 102 136	nap Score
250	281 159 202	79 153 154 196 185 127	190 180 349 137	109 106 178	Score 432 770
2.80E-20	1.30E-24 2.50E-10 2.90E-15	0.00063 5.80E-10 1.80E-10 1.30E-15 4.00E-14 3.40E-06	1.50E-13 2.20E-13 3.30E-31 2.10E-08	4.90E-06 4.50E-06 6.20E-13	Prob 1.30E-40 2.60E-23
43	73 38 47	43 36 35 31	35 38 62 53	38 40 4 38 60 41	% Ident C 56 54
20	59 10	13 14 23 95 26	11 18 14		Cvrg : 36 : 37
AMINOTRANSFERASE (ISOMERIZING) (HEXOSEPHOSPHATE AMINOTRANSFERASE) (D-FRUCTOSE-6-PHOSPHATE AMIDOTRANSFERASE) (GFAT) (L-GLUTAMINE-D-FRUCTOSE-6-PHOSPHATE AMIDOTRANSFERASE) (GLUCOSAMINE-6-PH [Bacillus subtilis] HYPOTHETICAL 80.1 KD PROTEIN IN SODA-COMGA	50S RIBOSOMAL PROTEIN L11 [Bacillus stearothermophilus, Peptide, 133 aa] (U93874) formate dehydrogenase chain A [Bacillus subtilis] GLUCOSAMINEFRUCTOSE-6-PHOSPHATE	INTERGENIC REGION [Bacillus subtilis] DNA PRIMASE [Bacillus subtilis] DNA PRIMASE [Listeria monocytogenes] RECF PROTEIN [Bacillus subtilis] (AF084104) hypothetical protein [Bacillus firmus] (Z99105) similar to hypothetical proteins [Bacillus subtilis] GRAMICIDIN S SYNTHETASE II (GRAMICIDIN S BIOSYNTHESIS GRSB PROTEIN) [Brevibacillus brevis]	DCUB [Haemophilus influenzae Rd] ATP-DEPENDENT NUCLEASE SUBUNIT B [Bacillus subtilis] subtilis] carbamoyl-phosphate synthase, small chain (EC 6.3) - Methanococcus jannaschii [] GTP-BINDING PROTEIN LEPA [Bacillus subtilis] HYPOTHETICAL 56.1 KD PROTEIN IN MFD-DIVIC	[Bacillus subtilis] [Bacillus subtilis] HYPOTHETICAL 25.6 KD PROTEIN IN PLCB-LDH INTERGENIC REGION (ORFA) [Listeria monocytogenes] (M18327) LacOPZ-alpha peptide from pUC9; putative [cloning vectors] ANAEROBIC C4-DICARBOXYLATE TRANSPORTER	NCBI gi description subtilis] (AF027868) putative transporter [Bacillus subtilis] (790100) similar to 3-excacult acyl-carrier protein synthase

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1657 BEIGG3850 BEIG4059 1	Bt1G4058			1654 Bt1Gc3830 Bt1G4055 27	1653 Bt1Gc3826 Bt1G4054 1-2	·		1652 Bt1Gc3822 Bt1G4053 44		1651 Bt1Gc3821 Bt1G4052 59-	1650 Bt1Gc3819 Bt1G4051 26	1649 Bt1Gc3812 Bt1G4050 31		1648 Bt1Gc3817 Bt1G4049 29	1647 Bt1Gc3798 Bt1G4047 26		1646 Bt1Gc3796 Bt1G4046 1-2	1645 Bt1Gc3785 Bt1G4045 1-2		1644 Bt1Gc3779 Bt1G4044 1-5		1643 Bt1Gc3772 Bt1G4043 41		1642 Bt1Gc3760 Bt1G4042 500		1641 Bt1Gc3761 Bt1G4041 1-	1640 Bt1Gc3759 Bt1G4040 14	1639 Bt1Gc3747 Bt1G4039 1-1		d	SEQ Contig Id Gene Id Posi	
1-344 8				270-1	1-259			445-1		59-294 8	267-1	319-1		290-1	262-1		1-275	1-272		1-556 ։ չ		411-1 8		500-177 €		1-427	141-1	1-135			Position	
<u>g</u> 321919	g2293156	g1945051		g3025180	g4584148			g1172699		g1730925	g3345481	g994736) 	g4584150	g2635763		g732376	g629038		g127481		g1078767		g2501575		g4558012	g1708641	g3290175		q	NCBI 9i	
184	183	157		100	210			376		178	152	78	}	269	153		173	141		203	-	238		121		178	103	115		Score		•
79	237	229		96	225			378		186	190	91	?	285	213		206	126		259		266		140		173	129	135	•	Score	BlastP	
0.004	5.80E-20	9.00E-19		0.00048	6.70E-18			1.30E-34		1.50E-14	5.40E-14	0.00017		4.80E-25	1.10E-16		1.10E-16	3.50E-07		2.70E-22		7.60E-22		4.00E-09		1.60E-12	7.10E-08	5.40E-08		Prob	BlastP-	1
49		35		•	54			54		43	4	54	•	. 59	4		47	48		32		37		26 .		36	4 8	61		Ident Cvrg	%	
77	91	43		20	9			26		45	15	57	}	31	18		36	17		97		16		35			15	7		Vrg	%	
hypothetical 16.9K protein - Salmonella typhimurium plasmid NTP16 []	(AF008220) YtiB [Bacillus subtilis]	(U63928) L1 protein [Bacillus cereus]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 49.9 KD PROTEIN IN CITA-SSPB	(AJ010111) pyruvate carboxylase [Bacillus cereus]	SYSTEM, ENZYME I) [Bacillus stearothermophilus]	PHOSPHOTRANSFERASE (PHOSPHOTRANSFERASE	PHOSPHOENOLPYRUVATE-PROTEIN	INTERGENIC REGION [Bacillus subtilis]	SP.J HYPOTHETICAL 19.0 KD PROTEIN IN ILVD-THYB	(AB016285) phosphotransferase system enzyme I [Bacillus	(M18327) LacOPZ-alpha peptide from pUC9; putative	[Bacillus cereus]	(AJ010111) cytochrome caa3 oxidase assembly factor	(Z99120) similar to hypothetical proteins [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 29.5 KD PROTEIN IN ROCC-PTA	hypothetical protein - Bacillus subtilis []	subtilis]	[Entamoeda histolytica] GTP CYCLOHYDROLASE I (GTP-CH-I) [Bacillus	alcohol dehydrogenase (EC 1.1.1.1) - Entamoeba histolytica	acetaldehyde dehydrogenase (acetylating) (EC 1.2.1.10)/	[Synechocystis sp.]	HYPOTHETICAL 33.3 KD PROTEIN SLL1263	Inhibitor []	Chain A, Alanine Racemase With Bound Propionate	(U51115) YeaC [Bacillus subtilis]	(AF067645) spore germination protein GerIA [Bacillus	INTERGENIC REGION [Bacillus subtilis]	9	NCBI of description	

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TRIGGER FACTOR (TF) (VEGETATIVE PROTEIN 2)	. 28	1 40	9.80E-14	185	168	g2829689	1-357	08 Bt1G4083	1680 Bt1Gc4008
(AF007865) bacitracin synthetase 3; BacC [Bacillus	_) 52	2.80E-10	167	165	g2982196	1-187	98 Bt1G4082	1679 Bt1Gc3998
HYPOTHETICAL 29.8 KD PROTEIN IN FRR-CDSA	. 25	7 42	9.20E-07	117	113	g3183468	282-88	83 Bt1G4081	1678 Bt1Gc3983
subtilis] ELONGATION FACTOR TS (EF-TS) [Bacillus subtilis]	35	1 39	9.20E-11	154	117	g3123214	1-306	82 Bt1G4080	1677 Bt1Gc3982
subtilis] (Z99124) pyrimidine-nucleoside transport protein [Bacillus	47	4 67	4.80E-64	653	651	g2636487	1-559	81 Bt1G4079	1676 Bt1Gc3981
subtilis] 30S RIBOSOMAL PROTEIN S11 (BS11) [Bacillus	97) 73	1.50E-30	337	440	g133719	1-385	78 Bt1G4078	1675 Bt1Gc3978
ORIGIN REGION [] (Z99110) similar to cystathionine beta-lyase [Bacillus	21	8 36	8.10E-08	130	99	g2633542	251-1	74 Bt1G4077	1674 Bt1Gc3974
[Enterobacteriaceae] HYPOTHETICAL 6.8 KD PROTEIN IN REPLICATION	. 79	8 82	4.70E-18	219	201	g141263	403-252	70 Bt1G4076	1673 Bt1Gc3970
INTERGENIC REGION [Bacillus subtilis] TRANSCRIPTIONAL REPRESSOR PROTEIN KORB	22	7 98	3.10E-37	400	402	g125524	238-1	69 Bt1G4075	1672 Bt1Gc3969
HYPOTHETICAL 61.5 KD PROTEIN IN ADEC-PDHA	26	5 34	1.50E-15	204	155	g2833392	435-1	66 Bt1G4074	1671 Bt1Gc3966
GCPE PROTEIN HOMOLOG [Bacillus subtilis]	23	2 45	2.50E-12	171	153	g1730252	1-263	64 Bt1G4073	1670 Bt1Gc3964
orthinine aminotransferase rocD - Bacillus subtilis []	25	3 42	3.00E-13	180	136	g2117766	301-1	60 Bt1G4072	1669 Bt1Gc3960
(X92970) orfB [Escherichia coli]	61	+ 33	8.20E-14	179	161	g1064900	445-1	59 Bt1G4071	1668 Bt1Gc3959
(299120) similar to arsenate reductase [Bacillus subtilis]	92	66	5.80E-36	388	374	g2635777	330-1	53 Bt1G4070	1667 Bt1Gc3953
(U89796) chitinase [Bacillus thuringiensis]	11	3 51	1.00E-13	188	162	g2149596	1-206	17 Bt1G4069	1666 Bt1Gc3917
HOMOLOG. [Bacillus subtilis]						(
(AB001488) ATP-DEPENDENT RNA HELICASE DEAD		4 8	3.80E-23	273	263	g1881268	1-407		
(AJ010131) vfkH [Bacillus cereus]		_	6.90E-17	208	224	g4584097	310-1	29 Bt1G4067	1664 Bt1Gc3929
acia nydrolase (pcbb) [Archaeoglobus rugidus] HIT PROTEIN [Bacillus subtilis]	32	-	4.30E-08	125	101	g3023940	1-140	00 Bt1G4066	1663 Bt1Gc3900
(AE000986) 2-hydroxy-6-oxo-6-phenylhexa-2,4-dienoic	56	7 40	8.80E-17	207	163	g2648849	79-490	70 Bt1G4065	1662 Bt1Gc3870
LINE-1 REVERSE TRANSCRIPTASE HOMOLOG []			9.50E-29	333	240	g126295	1-491		
			9.30E-12	166	152	g1945051	1-317		
subtilis]					•				•
subtilis] ACETOIN UTILIZATION ACUC PROTEIN [Bacillus	29	- - 4	9.20E-11	157	174	g728801	1-341	55 Bt1G4061	1659 Bt1Gc3855
THIAMIN BIOSYNTHESIS PROTEIN THIC [Bacillus	15	2	1.60E-22	269	259	g3041750	1-266	52 Bt1G4060	1658 Bt1Gc3852
NCBI gi description	Cyrg	70 Ident	Prob	Score	nap Score	NCBI gi	Position	ld Gene Id	ID Contig Id
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ACETYL-COENZYME A SYNTHETASE (ACETATE COA LIGASE) (ACYL-ACTIVATING ENZYME)	24 A	51	8.30E-32	353	303	g728788	1-410	Bt1G4103	1699 Bt1Gc4146 Bt1G4103	16
IN LEKGENIC REGION [Bacillus subtilis] TRANSPOSASE FOR INSERTION SEQUENCE	24 T	4	1.50E-17	220	177	g2497382	304-1	Bt1G4102	1698 Bt1Gc4135	16 -
(1 14080) nypometicai protein [Bacillus Suotilis] HYPOTHETICAL 30.7 KD PROTEIN IN MCPC-KINA	36 H	48	8.80E-17	207	208	g3183454	501-187	Bt1G4101	1697 Bt1Gc4136	16
(ORF73) [Bacillus subtilis]		ີ່ວ	0 005 13	170	175	G2226168	1_317	B+1C4100		-
PROBABLE ABC TRANSPORTER PERMEASE PROTEIN IN SODA-COMGA INTERGENIC REGION	34 P	34	9.60E-07	118	112	g1177035	299-1	Bt1G4099	1695 Bt1Gc4097	16
KINASE) (TH KINASE) [Bacillus subtilis]	× t	-		,					· ·	
HYDROXYETHYLTHIAZOLE KINASE (4-METHYL-5-	50 H	31	7.40E-10	. 145	116	g732340	407-1	Bt1G4098	1694 Bt1Gc4091	16
STAGE V SPORULATION PROTEIN S [Bacillus subtilis]	98 S	56	4.80E-09	134	195	g1174408	360-108	Bt1G4097	1693 Bt1Gc4090	16
typnimurium] 50S RIBOSOMAL PROTEIN L6 (BL10) [Bacillus subtilis]	32 S	73	1.20E-19	234	237	g1173058	1-175	Bt1G4096	1692 Bt1Gc4077	16
MG(2+) TRANSPORT ATPASE, P-TYPE 1 [Salmonella	12 M	33	3.40E-15	204	160	g543864	348-1	Bt1G4095	1691 Bt1Gc4075	16
PROTEIN 15) (VEG15) [Bacillus subtilis]		,		·.,	·					
PHOSPHORYLASE) (PNPASE) (VEGETATIVE	<u> </u>									
POLYRIBONUCLEOTIDE	10 P	63	2.80E-14	194	177	g1709698	1-203	Bt1G4094	1690 Bt1Gc4060	16
(Y16739) formate C-acetyltransferase [Piromyces sp.]		43	1.60E-23	281	230	g4218500	446-1	Bt1G4093	1689 Bt1Gc4055	16
(Z99112) mutS [Bacillus subtilis]	_	37	6.80E-18	229	223	g2634076	1-419	Bt1G4091	1688 Bt1Gc4040	16
(Y17332) proline-rich protein [Zea mays]	25 ()	27	. 0.42	49	95	g4138732	1-284	Bt1G4090	1687 Bt1Gc4039	16
HISF PROTEIN (CYCLASE) [Bacillus subtilis]	48 H	39	7.80E-05	63	162	g3913859	108-476	Bt1G4089	1686 Bt1Gc4033	16
(AB011836) similar to B. subtilis yerN gene(87% identity)	18 27 T	58	1.10E-15	204	201	g4512347	1-254	Bt1G4088	1685 Bt1Gc4027	16
[Bacillus subtilis]					}.					
(Z99112) chromosome segregation SMC protein homolg		32	7.70E-13	177	170	g2633966	1-501	Bt1G4087	1684 Bt1Gc4013	16
PRKA PROTEIN [Bacillus subtilis]	25 PI	36	4.40E-16	210	159	g730399	464-1	Bt1G4086	1683 Bt1Gc4023	16
Bacillus subtilis		i		į	`			(
[Bacillus subtilis] (Z99112) signal recognition particle (docking protein)	32 (F	32	7.30E-07	120	9 .	e2633967	316-1	Bt1G4085	1682 Bt1Gc4011	. 16
(vEGZ) [Bacillus subtilis respiratory nitrate reductase alpha chain -;Bacillus subtilis	6 re	53	8.20E-12	174	148	g2117582	227-1	Bt1G4084	1681 Bt1Gc4006	Ì6
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NCBI gi description	Vrg	% % Ident Cvrg	BlastP- Prob	BlastP Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	N E S
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Table 1

23 HYPOTHETICAL HELICASE IN SINI-GCVT INTERGENIC REGION [Bacillus subtilis]	35	0.069	77	142	g1731040	380-1	1716 Bt1Gc4203 Bt1G4120	171
thermophilus] 60 HYPOTHETICAL PROTEIN UL127 [human herpesvirus 5]	90	3.20E-35	381	367	g137018	1-235	5 Bt1Gc4201 Bt1G4119	1715
	52	1.80E-07	132	114	g2492564	164-1	Bt1Gc4193	1714
TRANSFER SUBUNIT [Bacillus subtilis] 30 (D88121) CPRD12 protein [Vigna unguiculata]	45	7.90E-13	170	131	g1853968	1-254	3 Bt1Gc4200 Bt1G4117	1713
[Bacillus subtilis] 27 DIHYDROOROTATE DEHYDROGENASE ELECTRON	36	3.20E-06	112	97	g141329	207-1	2 Bt1Gc4198 Bt1G4116	1712
11 EXCINUCLEASE ABC SUBUNIT B (DINA PROTEIN)	40	1.20E-08	141	112	g3123297	1-213	1 Bt1Gc4194 Bt1G4115	1711
PROTEIN 31) (VEG31) [Bacillus subtilis] 14 (Z99120) similar to hypothetical proteins [Bacillus subtilis]	69	7.60E-10	150	132	g2635763	197-1	0 Bt1Gc4188 Bt1G4114	1710
27 ATP SYNTHASE BETA CHAIN (VEGETATIVE	43	2.20E-14	192	218	g584808	378-1	9 Bt1Gc4179 Bt1G4113	1709
(TRANSCRIPTASE BETA CHAIN) (RNA POLYMERASE BETA SUBUNIT) [Bacillus subtilis]								
6 DNA-DIRECTED RNA POLYMERASE BETA CHAIN	46	1.10E-07	135	139	ģ585920	1-204	1708 Bt1Gc4177 Bt1G4112	170
16 TOPOISOMERASE IV SUBUNIT B [Bacillus subtilis]	41	2.50E-14	194	163	g3914289	313-1	7 Bt1Gc4174 Bt1G4111	1707
INTERGENIC REGION [Bacillus subtilis]								
35 HYPOTHETICAL 51.5 KD PROTEIN IN CITA-SSPB	42 .	1.30E-15	203	252	g3025181	539-41	6 Bt1Gc4172 Bt1G4110	1706
7 ISOLEUCYL-TRNA SYNTHETASE (ISOLEUCINE	 48	2.10E-09	150	98	g3123287	189-1	5 Bt1Gc4160 Bt1G4109	1705
	;		,	,				į
(IMIDAZOLONEPROPIONATE HYDROLASE)								-
formyltransferase [Bacillus cereus] 13 UROCANATE HYDRATASE (UROCANASE)	51	9.50E-09	141	94	g123765	242-459	1704 Bt1Gc4154 Bt1G4108	170
29 (Y09252) phosphoribosylaminoimidazolecarboxamide	35	1.00E-08	·137	107	g2072373	293-1	3 Bt1Gc4153 Bt1G4107	1703
(CARBAMOTIC-FROSFRATE STRIFETASE AMMONIA CHAIN) [Bacillus caldolyticus]					-			
PYRIMIDINE-SPECIFIC, LARGE CHAIN			•					
9 CARBAMOYL-PHOSPHATE SYNTHASE,	6 .	9.10E-15	201	146	g1168772	1-299	1702 Bt1Gc4144 Bt1G4106	170
subtilisi	Ņ	2.90E-08	136	105	g121516	1-248	1 Bt1Gc4150 Bt1G4105	10/1
	39	6.30E-07	114	106	g144312	123-395	Bt1Gc4142	1700
(ACETYL-COA SYNTHASE) [Bacillus subtilis]	•		•				-	
	% % Ident Cvrg	BlastP- Prob	BlastP Score	aat_ nap Score	NCBI gi	Position	Contig Id Gene Id	NO SEQ
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	1735 Bt1Gc4351 Bt1G4140	1733 Bt1Gc4340 Bt1G4138 1734 Bt1Gc4341 Bt1G4139	1732 Bt1Gc4337 Bt1G4137	1730 Bt1Gc4312 Bt1G4135	Bt1Gc4281	1/2/ Bt1Gc4263 Bt1G4132 1728 Bt1Gc4301 Bt1G4133		1725 Bt1Gc4264 Bt1G4130	1723 Bt1Gc4255 Bt1G4128 1724 Bt1Gc4253 Bt1G4129	1722 Bt1Gc4254 Bt1G4127	1720 Bt1Gc4241 Bt1G4124 1721 Bt1Gc4240 Bt1G4125	1719 Bt1Gc4238 Bt1G4123		1717 Bt1Gc4232 Bt1G4121	SEQ ID Contig Id Gene Id
	0 1-219	8 190-319 9 1-287	7 237-1	5 107-197 6 1-229		2 242-1 3 1-314		0 187-1	8 79-248 9 1-235	7 280-1	4 231-29 5 1-253	3 1-288		1 249-1	Position
es te	g2500053	g2633489 g585481	g1172465	g3183561 g2239276	g135811	g41/332 g3290177	g2635778	g2497382	g3915204 g595768	g2497392	g2497856 g133466	g2462959	g3122350	g1805393	NCBI gi
	121	75 252	320	90 386	153	133	154	115	123 111	181	116 99	191	113	127	aat_ nap
	148	1115 212	322	108 399	200	175	180	159	160 118	199	131 114	177	130	147	BlastP Score
	1.20E-09	1.60E-06 6.90E-16	5.70E-29	9.30E-06 4.00E-37	1.20E-15	7.90E-13	6.70E-13	6.90E-11	1.90E-10 2.40E-07	6.20E-16	4.00E-08 4.10E-06	1.30E-13	3.50E-08	1.90E-09	BlastP- Prob
	45	41 57	_	95		3 ¥ 8 4		52	61	49	41 33	3 %		42	% Ident C
	16	. 18 8	40	13	34	30	20	14	47	37	21 23	48	23	16	% Cyrg
	(1RCF) [Bacillus subtilis] 6 DNA REPAIR PROTEIN RADA HOMOLOG (DNA REPAIR PROTEIN SMS HOMOLOG) [Listeria			 VEGETATIVE PROTEIN 296 (VEG296) [Bacillus subtilis]. (L77763) neutral protease A [Bacillus thuringiensis] 			- ·	vector] 4 TRANSPOSASE FOR INSERTION SEQUENCE ELEMENT IS232 [Insertion sequence IS232]		A) (SIGMA-43) [Bacillus subtilis] 7 INSERTION SEQUENCE IS232 PUTATIVE ATP- BINDING PROTEIN [Insertion sequence IS232]	[Bacillus Subtilis] 1 MALATE DEHYDROGENASE [Bacillus israeli] 3 RNA POLYMERASE SIGMA FACTOR RPOD (SIGMA-		dehyd PROL	16 (D50453) 68% identity protein to 1-pyrroline-5-carboxylate	NCBI gi description

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474-217	1751 Bt1Gc4441 Bt1G4157 1-302 g2632593	1750 Bt1Gc4436 Bt1G4156 1-367 g2293213	1749 Bt1Gc4432 Bt1G4155 286-1 g2635778		1748 Bt1Gc4416 Bt1G4154 1-102 g3892204	•	1747 Bt1Gc4407 Bt1G4153 132-1 g1711644				1746 Bt1Gc4391 Bt1G4152 388-28 g584771		1745 Bt1Gc4384 Bt1G4151 1-389 g2634887	1744 Bt1Gc4382 Bt1G4150 345-32 g1934806		1743 Bt1Gc4381 Bt1G4149 1-333 g1168649	-	1742 Bt1Gc4370 Bt1G4148 320-1 g1881268		1741 Bt1Gc4367 Bt1G4147 1-311 g3777577	,	1740 Bt1Gc4368 Bt1G4146 1-403 g267497	•	1739 Bt1Gc4360 Bt1G4145 379-1 g2497382	•	1738 Bt1Gc4358 Bt1G4144 1-264 g3329623	1/3/ BII/G4334 BII/G4143 309-1 g2829689		1736 Bt1Gc4355 Bt1G4141 1-278 g1168646		Contig Id Gene Id Position NCBI gi	
145	175	479	67		154		115		-		91		198	104		182		129		184		137		270		108	221	3	133		ю .	88t R
152	112	483	116		105		134				112		217	140		213		117		110		186		276		66	241	:	132		Score	Get D
2.40E-10	8.50E-06	5.00E-46	5.10E-06		0.00015		4.80E-08				8.00E-06		7.70E-18	1.20E-09		2.00E-17		3.20E-06		6.10E-06		9.30E-14		7.20E-24	,	0.43	6.40E-20		2.90E-08		Prob	Riest D
36	41	~75	37		94		62				30		40	39		43		36		47		33		49		27	47	·	41		Ident Cyrg	,
25	23	61	17		ω		9				29		46	47		4		· 21		38		. 26		29		26	24		31		Cyrg	\$
NARA PROTEIN [Bacillus subtilis]	(Z99105) multidrug-efflux transporter [Bacillus subtilis]	(AF008220) YtpR [Bacillus subtilis]	(Z99120) similar to butyryl-CoA dehydrogenase [Bacillus	binding proteins [Cricetulus griseus]	[Synechocystis sp.] (AF078105) site-1 protease of sterol regulatory element	(ASPARAGINETRNA LIGASE) (ASNRS)	PROBABLE ASPARAGINYL-TRNA SYNTHETASE	SYNTHASE) (AGS) [Bacillus stearothermonhilus]	TRANSACETYLASE) (OATASE) / AMINO-ACID	ACETYLTRANSFERASE) (ORNITHINE	GLUTAMATE N-ACETYLTRANSFERASE (ORNITHINE	[Bacillus subtilis]	(Z99116) similar to hypothetical proteins from B. subtilis	(Z93936) unknown [Bacillus subtilis]	CYTOCHROME B/C SUBUNIT [Bacillus subtilis]	MENAQUINOL-CYTOCHROME C REDUCTASE	HOMOLOG. [Bacillus subtilis]	(AB001488) ATP-DEPENDENT RNA HELICASE DEAD	vector p34S-Km3]	(AF083408) kanamycin 3'-phosphotransferase [Cloning	3'REGION (ORF A) [Bacillus firmus]	HYPOTHETICAL SO.9 KD PROTEIN IN KATA	ELEMENT IS232 [Insertion sequence IS232]	TRANSPOSASE FOR INSERTION SEQUENCE	elegans	(VEG2) [Bacillus subtilis] (AF078790) No definition line found [Caenorhabditis]	IRIGGER FACTOR (IF) (VEGETATIVE PROTEIN 2)	PROTEIN) [Bacillus subtilis]	GTP-BINDING PROTEIN ERA HOMOLOG (BEX	monocytogenes]	NCBI gi description	

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(AF008220) SAM synthase [Bacillus subtilis]	60 20	1.80E-19	236	231	g2293164	1-240	1793 Bt1Gc4750 Bt1G4200	
CATALASE []		6.30E-08	132	125	g231698	1-287	Bt1Gc4746	
NODULIN 21 (N-21) [Glycine max]		3.30E-10	145	102	g128405	1-373	Bt1Gc4737	
(Z75208) hypothetical protein [Bacillus subtilis]	50 35	5.40E-17	209	191	g1770029	1-270	1790 Bt1Gc4719 Bt1G4197	
(AJ224964) preprotein translocase [Phormidium	34 21	3.00E-06	118	. 113	83288383	4-3/1	1/89 BttGc4/14 BttG4196	
(U138/1) lacz alpna peptide [Cioning vector]		1.90E-06	110	122	2227507 08/ C6CB	323-220	B11Gc4/03	
STAGE V SPORULATION PROTEIN I [Bacillus supriis]		4.10E-12	100	100	g280883	148-1	Bt1Gc4701	
(D8/9/9) YmJ [Bacillus subtilis]		2.00E-23	282	150	g21169/4	433-1	Bt1 Gc4690	
(AF007865) Back [Bacillus licheniformis]	٠	6.70E-19	227	206	g4481749	291-1	Bt1Gc4695	
subtilis]	•		})		
IRON-UPTAKE SYSTEM PROTEIN FEUB [Bacillus	39 36	9.00E-18	217	139	g729484	377-18	1784 Bt1Gc4691 Bt1G4191	
PYRUVATE KINASE (PK) [Synechocystis sp.]	32 33	3.10E-15	200	181	g2497534	472-1	1783 Bt1Gc4687 Bt1G4190	
influenzae GB:U00019_14 [Bacillus cereus]			,					
(Y11140) homology to M. leprae B2235_C2_195 and H.	72 37	1.20E-28	319	317	g2462120	1-253	1782 Bt1Gc4681 Bt1G4189	
SYNTHETASE) [Bacillus subtilis]								
(GLUTAMINE AMIDOTRANSFERASE) (GMP								
GMP SYNTHASE (GLUTAMINE-HYDROLYZING)	56 27	8.30E-37	396	379	g3123227	1-429	1781 Bt1Gc4679 Bt1G4188	
[Bacillus subtilis]			•					
(AF008553) Glu-tRNAGln amidotransferase subunit A	60 18	4.90E-20	244	202	g2589195	271-1	1780 Bt1Gc4673 Bt1G4187	
INTERGENIC REGION [Bacillus subtilis]								
HYPOTHETICAL 13.0 KD PROTEIN IN QCRC-DAPB	48 75	1.20E-14	187	152	g1176702	1-258	1779 Bt1Gc4674 Bt1G4186	
(ACONITASE) [Bacillus subtilis]			,					
ACONITATE HYDRATASE (CITRATE HYDRO-LYASE)	45 12	1.00E-15	209	171	g2506131	1-334	1778 Bt1Gc4670 Bt1G4185	
(D88802) transmembrane [Bacillus subtilis]	40 62	2.40E-23	269	287	g1945117	1-459	1777 Bt1Gc4659 Bt1G4184	
INTERGENIC REGION [Bacillus subtilis]								
[Pseudomonas aeruginosa] HYPOTHETICAL 36.3 KD PROTEIN IN RECQ-CMK	48 25	2.90E-15	195	165	g1730898	1-247	1776 Bt1Gc4657 Bt1G4183	
DEHYDROGENASE (ACYLATING) (MMSDH)								
METHYLMALONATE-SEMIALDEHYDE	44 12	0.0085	85	107	g127210	1-186	1775 Bt1Gc4644 Bt1G4182	
(299118) similar to single-suand DNA-specific exonuclease	41 12	1.00E-09	132	124	82033220	1-200	1//4 500004010 5004181	
(AB007638) function unknown [Bacillus subtilis]		1.20E-10	149	112	g2522020	125-337	Bt1Gc4598	
(U93876) hypothetical protein YrdR [Bacillus subtilis]		6.50E-05	102	80	g1934659	1-148	Bt1Gc4596	
stearothermophilus]								
NCBI gi description	. 🚡	·	Score	nap Score	NCBI gi	Position	ID Contig Id Gene Id NO	
	% %	RiastP. (RiastP	aat			SEQ	7.0

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-	37 29	2.10E-14	192	196	g2635847	406-1	1812 Bt1Gc4876 Bt1G4219	
DEHYDROGENASE (NADP+)) (IRIOSEPHOSPHAIE DEHYDROGENASE) [Zea mays] 3 N-ACETYL-GAMMA-GLUTAMYL-PHOSPHATE REDIICTASE (AGPR) (N-ACETYL-GLIITAMATE	47 23	3.40E-07	122	145	g584768	205-2	1811 Bt1Gc4862 Bt1G4218	
PHOSPHATE DEHYDROGENASE) (GLYCERALDEHYDE-3-PHOSPHATE	•		Ą	. 5				
		5.50E-11	161	126	g2494075	1-415	Bt1Gc4869	
4 ATP SYNTHASE AI PHA CHAIN [Bacillus megaterium]	46 14	6.30E-09	142	124	g114510	1-213	1809 Bt1Gc4856 Bt1G4216	
(Z939		8.60E-09	142	119	g1934835	1-259	Bt1Gc4851	
[Bacillus subtilis]			;					
	42 41	2.50E-28	. 316	322	g1705442	1-449	1806 Bt1Gc4809 Bt1G4213	
6 (X98455) SNF [Bacillus cereus]	47	2.70E-07	131	109	g1769947	228-39	1805 Bt1Gc4808 Bt1G4212	
	٠				. (
carboxyvinyirransierase [Bacillus subilis] 2 TRANSPOSASE FOR INSERTION SEQUENCE	56 22	1.10E-20	248	213	g2497382	1-287	1804 Bt1Gc4802 Bt1G4211	
	44 · 25	9.20E-18	222	199	g1648861	155-480	1803 Bt1Gc4804 Bt1G4210	
pallidum]								
	29 26	3.50E-11	164	134	g3323354	1-460	1802 Bt1Gc4798 Bt1G4209	
_	76 43	6.30E-23	265	246	g2340005	1-208	1801 Bt1Gc4795 Bt1G4208	
AMINOTRANSFERASE) (GSA-AT) [Bacillus subtilis]				٠.				
GLUI	58 36	2.90E-34	372	372	g399784	1-470	1800 Bt1Gc4791 Bt1G4207	
crypto	37. 10	2.30E-05	110	80	g2129564	185-1	1799 Bt1Gc4781 Bt1G4206	
[Bacillus stearothermophilus]					1			
		9.60E-13	182	131	g2661433	453-1	1798 Bt1Gc4779 Bt1G4205	
	51 18	3.00E-13	180	160	g1196998	254-31	1797 Bt1Gc4773 Bt1G4204	
		2.40E-05	111	72	g2633696	336-87	1796 Bt1Gc4763 Bt1G4203	
	-	2.50E-16	208	220	g1196998	131-423		
8 (AF008220) YtmQ [Bacillus subtilis]	36 38	9.70E-08	123	110	g2293207	1-249	1794 Bt1Gc4730 Bt1G4201	
NCBI gi description	¥ .		Score	nap Score	NCBI 81	Position	NO Contigua Genera	
	%	BlastP-	BlastP	aat_ j				

Table 1

				~																													
1833 Btl	1832 Bt1	1831 Bt1	1830 Bt1	1829 Bt1	1828 Bt1	1827 Bt1		1826 Bt1	1825 Bt1		1824 Bt1Gc4956	. <		1822 Bt1	1821 Bt1		-		_			1820 Bt1	1819 Bt1	1818 Bt1	1817 Bt1		1816 Bt1	1815 Bt1	1814 Btl	1813 Bt1		O	E Co
Bt1Gc5039	Bt1Gc5027	Bt1Gc5021	Bt1Gc5005	Bt1Gc5003	Bt1Gc5004	Bt1Gc5002		Bt1Gc4997	Bt1Gc4962		Gc4956			Bt1Gc4921	Bt1Gc4920							Bt1Gc4914	Bt1Gc4905	Bt1Gc4903	Bt1Gc4895		Bt1Gc4894	Bt1Gc4890	Bt1Gc4884	Gc4881			Contig Id
Bt1G4240	Bt1G4239	Bt1G4238	Bt1G4237	Bt1G4236	Bt1G4235	Bt1G4234		Bt1G4233	Bt1G4232		Bt1G4231	٠٠.	Bt1G4230	Bt1G4229	Bt1G4228							Bt1G4227	Bt1G4226	Bt1G4225	Bt1G4224		Bt1G4223	Bt1G4222	Bt1G4221	Bt1Gc4881 Bt1G4220			Gene Id
327-1	279-1	1-306	383-1	1-461	1-388	206-1		20-379	160-1		1-189		420-21	203-1	1-186					•		1-262	1-227	1-458	1-377		384-1	. 245-1	1-169	1-224			Position
g3123297	g732355	g2116974	g1945657	g2635858	g2635600	g595780		g3328184	g790866	•	g732384	(g2127273	g1124822	g2988346						÷.	g1169919	g2635181	g2116767	g2116767	9	g2501371	g4033397	g225559	g1695686			NCBI gi
203	107	224	138	207	98	104		. 225	95		96		679	182	112					`.	٠.	145	115	388	231	:	170	145	169	133		Score	nap_
233	135	263	160	259	128	92		235	113		106		681	212	156							149	202	332	273		184	183	173	115	•	0	BlastP
1.60E-18	8.20E-09	2.20E-21	1.10E-11	3.50E-21	1.30E-07	0.00025		9.50E-20	2.70E-06		2.70E-05		5.20E-67	1.50E-16	2.00E-10							1.50E-09	3.10E-16	3.00E-29	7.60E-23	!	2.40F-14	5.40E-13	2.60E-12	1.50E-05			BlastP-
54	33	54	33	34	19	47		43	46		40		98	66	49							3 8	60	52	40	•	36	43	63	48			Ident Cyro
16	36	10	49	20	33	37		42	20		18		&		12						•	15	24	23	19	;	4	10	12	6		ď	* %
INTERGENIC REGION [Bacillus subtilis] EXCINUCLEASE ABC SUBUNIT B (DINA PROTEIN) [Bacillus subtilis]	HYPOTHETICAL 28.4 KD PROTEIN IN SACT-SACP	(D87979) YfnJ [Bacillus subtilis]	(Z94043) hypothetical protein [Bacillus subtilis]	(Z99121) yvgS [Bacillus subtilis]	(Z99119) similar to hypothetical proteins [Bacillus subtilis]	(U13871) lacZ alpha peptide [Cloning vector]	pQE30NST]	(AF074376) beta lactamase precursor [Cloning vector	(L36381) orfB; putative [Neisseria gonorrhoeae]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 37.0 KD PROTEIN IN SPOIIR-GLYC		B. subtilis spoIIGA protein homolog - Bacillus thuringiensis	(M16158) unknown protein [Bacillus thuringiensis]	(D89812) catalase [Sus scrofa]	[Bacillus subtilis]	AMIDOTRANSFERASE) (GLUCOSAMINE-6-PH	(GFAT) (L-GLUTAMINE-D-FRUCTOSE-6-PHOSPHATE	FRUCTOSE-6-PHOSPHATE AMIDOTRANSFERASE)	(HEXOSEPHOSPHATE AMINOTRANSFERASE) (D-	AMINOTRANSFERASE (ISOMERIZING)	GLUCOSAMINEFRUCTOSE-6-PHOSPHATE	(Z99117) similar to protease [Bacillus subtilis]	(D86418) YfnI [Bacillus subtilis]		CYSW [Symechocyctic en]	SUIL FATE TRANSPORT SYSTEM PERMEASE PROTEIN	DNA GYRASE SUBUNIT A [Bacillus sp.]	stearothermophilus of ORF iS231C [Bacillus thuringiensis]	(D83706) pyruvate carboxylase [Bacillus	[Bacillus subtilis]		NCBI gi description

Table 1

94.7	1854 Bt		1852 Bt	1851 Bt		1850 Bt	1849 Bt		1848 Bt		1847 Bt	1846 Bt			1844 Bt			1843 Bt		1842 Bt	1841 Bt	1840 Bt	1839 Bt	1838 Bt	1837 Bt	1836 Bt	1835 Bt	1834 Bt	SEQ C
	Bt1Gc5183	Bt1Gc5171	Bt1Gc5166	Bt1Gc5162		Bt1Gc5161	Bt1Gc5159		Bt1Gc5150	. • * :	Bt1Gc5144	Bt1Gc5131	Bt1Gc5130) # 	1844 Bt1Gc5114	•		Bt1Gc5098		Bt1Gc5087	Bt1Gc5080	Bt1Gc5081	Bt1Gc5073	Bt1Gc5070	Bt1Gc5069	Bt1Gc5065	Bt1Gc5056	Bt1Gc5047	Contig Id
	Bt1G4261	Bt1G4260	Bt1G4259	Bt1G4258		Bt1G4257	Bt1G4256		Bt1G4255		Bt1G4254	Bt1G4253	Bt1G4252		Bt1G4251			Bt1G4250		Bt1G4249	Bt1G4248	Bt1G4247	Bt1G4246	Bt1G4245	Bt1G4244	Bt1G4243	Bt1G4242	Bt1G4241	Gene Id
	450-1	1-300	220-1	1-386		1-139	67-362		1-386		374-1	1-256	1-395		1-446			1-369	,	489-1	1-379	34-321	114-522	400-26	314-63	57-296	1-421	1-358	Position
	g1945096	g2443243	g2635763	g2462090	-	g2144427	g2226151	(g1176955		g585920	g1729789	g2635723		g1731364			g729340	0	g3025180	g730399	g266569	g2619012	g225559	g729341	g2577965	g2633541	g1619837	NCBI gi
	446	87	117	97		89	182		212		326	111	87		194			136		524	322	171	411	277	98	186	184	141	aat_ nap Score
	406	110	140	130		113	212		234		363	147	127		242	٠		220	į	524	344	150	436	219	138	197	240	67	BlastP Score
٠	7.20E-38	1.70E-06	9.20E-09	1.20E-07		1.30E-05	9.40E-17		2.90E-19		5.50E-32	1.50E-09	2.10E-07		3.70E-20			2.30E-17		2.30E-50	1.30E-30	2.50E-10	4.80E-41	2.60E-17	1.10E-08	1.40E-14	4.00E-20	0.00064	BlastP- Prob
	. 59	30	4	30		40	43		40		62	34	26		37			35	;	60	58.	41	61	48	34	48	40	39	% Ident
	35	85	16	25		7	24		32		10	19	30		37		-	24		36	20	32	39	. 26	21	11	37	39	%
	(D88802) S. livio P31141 (492) tra	(D86417) YfIT	(Z99120) similar	(Y11171) BC542A protein	glucose-specific,	phosphotransfera	(Y14080) hypoth	INTERGENIC R	HYPOTHETICA	(TRANSCRIPTA BETA SUBUNIT	aureus] DNA-DIRECTE	(Y09927) phosph	(Z99120) homos	INTERGENIC R	HYPOTHETICA	(DCL) [Bacillus subtilis]	ALANINE-D-AI	D-ALANINE-AC	INTERGENIC R	HYPOTHETICA	stearothermophilus] PRKA PROTEIN [ROD SHAPE-DI	(AF027868) fatty	ORF IS231C [Bi	DLTB PROTEIN	subtilis] (Y15254) PcrA p	(Z99110) similar	(U67921) collage	•
•	 S. lividans chloramphenicol resistance protein; (492) transmembrane [Bacillus subtilis] 	(D86417) YfIT [Bacillus subtilis]	to hypothetical prote	lococcus carnosus] 1) BC542A protein [Bacillus cereus]	-specific, factor IIB - Staphylococcus carnosus	phosphotransferase system enzyme II (EC 2.7.1.69),	(Y14080) hypothetical protein [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 45.8 KD PROTEIN IN ACDA-NARI	(TRANSCRIPTASE BETA CHAIN) BETA SUBUNIT) [Bacillus subtilis]	aureus] DNA-DIRECTED RNA POLYMERASE BETA CHAIN	7) phosphoglucosamine mutase [Staphylococcus	0) homoserine dehydrogenase [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL PROTEASE IN ROCR-PURA	subtilis]	ALANINE-D-ALANYL CARRIER PROTEIN LIGASE)	D-ALANINE-ACTIVATING ENZYME (DAE) (D-	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 49.9 KD PROTEIN IN CITA-SSPB	stearothermophilus] PRKA PROTEIN [Bacillus subtilis]	ETERMINING PROT	(AF027868) fatty acid desaturase [Bacillus subtilis]	ORF IS231C [Bacillus thuringiensis]	DLTB PROTEIN [Bacillus subtilis]	subtilis] (Y15254) PcrA protein [Bacillus subtilis]	to cystathionine gam	1) collagen-like protein [Bacillus thuringiensis	NCBI gi description
	l resistance protein; us subtilisl	-	similar to hypothetical proteins [Bacillus subtilis]	cereus]	coccus carnosus	(EC 2.7.1.69),	lus subtilis]	btilis]	N IN ACDA-NARI	(TRANSCRIPTASE BETA CHAIN) (RNA POLYMERASE BETA SUBUNIT) [Bacillus subtilis]	SE BETA CHAIN	e [Staphylococcus	[Bacillus subtilis]	btilis]	OCR-PURA		ROTEIN LIGASE)	VIE (DAE) (D-	htilie]	N IN CITA-SSPB		ROD SHAPE-DETERMINING PROTEIN MREC [Bacillus	cillus subtilis]		-	ilis]	israeiensis] (Z99110) similar to cystathionine gamma-synthase [Bacillus	llus thuringiensis	Ön
									•																				* *

Table 1

Table 1

	1896 Bt1Gc5563 Bt1G4306 1-449	1895 Bt1Gc5536 Bt1G4304 322-1	1894 Bt1Gc5527 Bt1G4303 306-1	1893 Bt1Gc5521 Bt1G4302 420-90	1892 Bt1Gc5501 Bt1G4301 600-1	1891 Bt1Gc5498 Bt1G4300 447-1	1890 Bt1Gc5488 Bt1G4299 465-1	1889 Bt1Gc5486 Bt1G4298 1-152		1888 Bt1Gc5467 Bt1G4297 1-255	1887 Bt1Gc5454 Bt1G4296 269-1	·	1886 Bt1Gc5473 Bt1G4295 278-1	1885 Bt1Gc5472 Bt1G4294 1-388		1884 Bt1Gc5466 Bt1G4293 1-144	1883 Bt1Gc5464 Bt1G4292 159-1		1882 Bt1Gc5453 Bt1G4291 596-295	1881 Bt1Gc5442 Bt1G4290 366-1	1880 Bt1Gc5412 Bt1G4289 388-1		18/9 BII GC3411 BII G4288 1-419		1878 Bt1Gc5406 Bt1G4287 382-1	1877 Bt1Gc5386 Bt1G4286 490-130	SEQ ID Contig Id Gene Id Position NO
			l g730776	0 g729325	1 g586848	1 g3123286							1 g2501426	8 g729583			1 g1009035		95 g2633170	1 g2635670	l g2618834		82492/3/		l g1731075	30 g2633738	on NCBI gi
	115	105	200	108	117	144	458	102		109	117		255	66	-	119	184		244	143	171		187		138	203	nap Score
	203	129	220	143	211	161	477	128		130	124		301	130		129	184		219	183	175	٠	228	3	151	203	BlastP Score
	1.40E-15	1.60E-08	5.40E-17	1.10E-09	7.90E-17	1.20E-10	2.20E-45	1.20E-07		7.80E-08	4.60E-07		9.70E-27	6.30E-08		1.60E-08	2.30E-13		6.50E-18	2.20E-13	1.60E-12		8.80E-18		3.00E-10	2.30E-16	BlastP- Prob
	42	36	49 .	32	34	31	63	53		3 8	32		62	21		60	62		47	33	28		3/	:	29	46	% Ident
	19	కు	13	42	53	18	32	13		22	20	•	21	. 33 . 00		24	9		29	22	27		16	;	38	. 86	%.
thermoautotrophicum]	PRISMANE PROTEIN HOMOLOG [Methanobacterium	(U66480) YnaD [Bacillus subțilis]	[Bacillus megaterium] STAGE III SPORULATION PROTEIN E [Bacillus subtilis]	RPSF-SP00J INTERGENIC REGION [Bacillus subtilis] GLUCOSE 1*DEHYDROGENASE II (GLCDH-II)	HYPOTHETICAL 40.1 KD GTP-BINDING PROTEIN IN	LEUCYL-TRNA SYNTHETASE (LEUCINETRNA	ORF IS231C [Bacillus thuringiensis]	(Z99112) nusA [Bacillus subtilis]	(ADP-GLUCUSE SYNTHASE) (ADP-GLUCUSE PYROPHOSPHORYLASE) [Bacillus subtilis]	GLUCOSE-1-PHOSPHATE ADENYLYLTRANSFERASE	(U63928) L2 protein [Bacillus cereus]	[Bacillus stearothermophilus]	PYRIMIDINE-NUCLEOSIDE PHOSPHORYLASE	GLYCOGEN BIOSYNTHESIS PROTEIN GLGD [Bacillus	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 21.4 KD PROTEIN IN DACA-SERS	(L47692) pyruvate oxidase [Escherichia coli]	[Bacillus subtilis]	[Bacillus subtilis] (Z99108) similar to iron(III) dicitrate transport permease	(Z99120) similar to nicotinate phosphoribosyltransferase	(AF017113) putative protease [Bacillus subtilis]	Entamoeha histolytical	ALCOHOL DEHYDROGENASE (ACDE)	OXIDOREDUCTASE YQJM [Bacillus subtilis]	PROBABLE NADH-DEPENDENT FLAVIN	(Z99111) similar to transcriptional regulator (MarR family)	NCBI gi description

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SEQ ID Contig Id Gene Id NO	Position	NCBI gi	aat_ j	BlastP Score	BlastP- Prob	% % Ident Cvrg	%	NCBI gi description
1897 Bt1Gc5568 Bt1G4307	356-1	g118793	302	292	2.70E-24	55	00	DNA POLYMERASE III, ALPHA CHAIN [Bacillus
1898 Bt1Gc5574 Bt1G4308	190-1	g3777577	127	146	5.60E-10	46	23	subtilis] (AF083408) kanamycin 3'-phosphotransferase [Cloning
1800 Bt1Gc5585 Bt1G4300	186-1	02462104	145	5	7 10E-11	ŝ	2	vector p34S-Km3]
	,	9	;	į		ţ	ţ	cereus]
1900 Bt1Gc5590 Bt1G4310	148-5	g121415	119	121	1.10E-06	46	9	GLYCEROL KINASE (ATP:GLYCEROL 3-
•								PHOSPHOTRANSFERASE) (GLYCEROKINASE) (GK) [Bacillus subtilis]
1901 Bt1Gc5592 Bt1G4311	1-188	g118672	125	, 13 8	1.50E-08	46	13	LIPOAMIDE DEHYDROGENASE COMPONENT OF PYRUVATE DEHYDROGENASE COMPLEX (E3)
						.,		(DIHYDROLIPOAMIDE DEHYDROGENASE) (S COMPLEX, 50 KD SUBUNIT) [Bacillus subtilis]
1902 Bt1Gc5609 Bt1G4312	1-496	g1789981	258	286	3.80E-25	35	58	(AE000433) IS150 putative transposase [Escherichia coli]
1903 Bt1Gc5627 Bt1G4313	171-1	g121887	92	114	9.20E-06	40	9	DNA GYRASE SUBUNIT B [Bacillus subtilis]
1904 Bt1Gc5637 Bt1G4314	1-150	g1709188	86	103	0.00014	. 43	00	DNA MISMATCH REPAIR PROTEIN MUTL [Bacillus subtilis]
1905 Bt1Gc5636 Bt1G4315	1-199	g2522016	85	106	2.70E-05	41	19	(AB007638) dehydrogenase [Bacillus subtilis]
1906 Bt1Gc5642 Bt1G4317	22-481	g115022	380	353	3.00E-32	58	. 59	BETA-LACTAMASE PRECURSOR, TYPE II (PENICILLINASE) (CEPHALOSPORINASE) [Bacillus
ا قم								cereus
1907 Bt1Gc5647 Bt1G4318	1-415	g117576	170	క	0.79	30	4	CIRCUMSPOROZOITE PROTEIN PRECURSOR (CS) [Plasmodium berghei]
Bt1Gc5656	1-409	g4584100	222	215	1.30E-17	43	81	(AJ010132) hypothetical protein [Bacillus cereus]
Bt1Gc5662	319-1	g4033396	300	321	4.40E-28	61	17	DNA GYRASE SUBUNIT B [Bacillus sp.]
1910 Bt1Gc5671 Bt1G4322	1-421	g3123300	275	261	1.70E-22	49	4	THIOREDOXIN REDUCTASE (GENERAL STRESS PROTEIN 35) (GSP35) [Bacillus subtilis]
Bt1Gc5671	1-421	g4379428	231	227	6.70E-19	46	45	(AJ223781) thioredoxin reductase [Staphylococcus aureus]
1911 Bt1Gc5675 Bt1G4324	161-1	g1524023	95	117	1.20E-06	49	19	(X95401) beta-lactamase [Escherichia coli]
1912 Bt1Gc5677 Bt1G4325	272-1	g2635723	153	8 3	0.0018	43	21	(Z99120) homoserine dehydrogenase [Bacillus subtilis]
1913 Bt1Gc5681 Bt1G4326	282-1	g1172710	158	170	4.90E-12	40	21	PTS SYSTEM, CELLOBIOSE-SPECIFIC IIC
		-					•	COMPONENT (EIIC-CEL) (CELLOBIOSE-PERMEASE
· .							•	ENZYME II, C COMPONENT) [Bacillus subtilis]
1914 Bt1Gc5686 Bt1G4327	1-275	g2668553	130	68	0.85	37	6	(U62929) multidrug resistance protein 1 [Filobasidiella
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Table 1

HEAT-INDUCIBLE TRANSCRIPTION REPRESSOR	24	51	2.40E-14	188	176	g3122229	1-244	Bt1G4347	1929 Bt1Gc5808 Bt1G4347
(TRANSCRIPTASE BETA CHAIN) (RNA POLYMERASE BETA SUBUNIT) [Staphylococcus aureus]									
DNA-DIRECTED RNA POLYMERASE BETA CHAIN	5	50	1.80E-07	133	117	g1350849	1-182	Bt1G4346	1928 Bt1Gc5802
(Z99120) similar to thioredoxin [Bacillus subtilis]	91	45	4.10E-19	229	198	g2635773	21-553	Bt1G4345	1927 Bt1Gc5796
(Z99120) yusF [Bacillus subtilis]	59	51	4.40E-15	191	228	g2635774	1-262	Bt1G4344	1927 Bt1Gc5796
[Bacillus subtilis]				•					
PEPTIDASE) (DD-CARBOXYPEPTIDASE) (PRP-5*)									
PENICILLIN-BINDING PROTEIN 5* PRECURSOR (D-	21	35	2.20E-08	135	77	g461914	263-1	Bt1G4342	1926 Bt1Gc5777
(X67138) flagellin [Bacillus thuringiensis]	55	52	3.90E-30	333	276	g1834395	1-459	_	
(M97678) Rho Factor [Bacillus subtilis]	25	59	1.30E-24	282	276	g143434	319-1	Bt1G4340	1924 Bt1Gc5772
(CATABOLITE CONTROL PROTEIN) [Bacillus subtilis]						,			
GLUCOSE-RESISTANCE AMYLASE REGULATOR	18	55	1.80E-10	153	133	g115950	254-70	Bt1G4339	1923 Bt1Gc5767
(ABOT1838) metyr-accepting chemotaxis protein (Bacillus halodurans)	3	ŧ	2.40E-13		331	84012307	1-500	סנלטאטסס	1322 Bt10c3/08
Subtilis]		.	2 12	106	3	2/612280	1 500		
ENOLPYRUVYL TRANSFERASE) (EPT) [Bacillus						٠			
TRANSFERASE) (UDP-N-ACETYLGLUCOSAMINE									
CARBOXYVINYLTRANSFERASE (ENOYLPYRUVATE									
PROBABLE UDP-N-ACETYLGLUCOSAMINE 1-		51	1.50E-30	337	256	g1171068	1-417	Bt1G4337	1921 Bt1Gc5771
ORF IS231C [Bacillus thuringiensis]	14	54	0.0035	· 55	153	g225559	1-205	Bt1G4336	1920 Bt1Gc5761
INTERGENIC REGION [Bacillus subtilis]									
HYPOTHETICAL 26.2 KD PROTEIN IN FTSH-CYSK	31	3 8	4.20E-09	136	131	g586893	212-1	Bt1G4335	1919 Bt1Gc5751
[Bacillus subtilis]								•	-
(SUPEROXIDE-INDUCIBLE PROTEIN 12) (SOU2)		•							
INOSINE-5'-MONOPHOSPHATE DEHYDROGENASE	27	37	0.0016	92	204	g124423	411-1	Bt1G4334	1918 Bt1Gc5748
sp.]						(
subuns] (AB015670) A2-5a orf21: hypothetical protein [Bacillus	45	40	1.10E-07	121	141	g3318590	1-240	Bt1G4333	1917 Bt1Gc5732
DIHYDRODIPICOLINATE REDUCTASE [Bacillus	32	53	9.80E-18	216	197	g1706300	380-125	Bt1G4332	1916 Bt1Gc5727
) [Bacillus halodurans]	;	. ;		i	!	9	 - -		
neoformans] (AB017508) truA homologue (identity of 62% to B. subtilis	35	28	3.40E-06	112	%	g4512436	282-1	Bt1G4330	1915 Bt1Gc5715
NCDI gi description	Cvrg	Ident	Prob	Score	nap Score	NCBI 81	TOSHIGHT	Clene In	NO Configura
NODI : Louistin	%	%	BlastP-	BlastP	aat_	NCBI 2:	Dogition	Canald	•
			Table 1			,			. <i>:</i>

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CHORISMATE SYNTHASE (5- ENOLPYRUVYLSHIKIMATE-3-PHOSPHATE	17	45	2.10E-08	135	108	g399058	186-1	Bt1G4370	1949 Bt1Gc5968	
ORF IS231C [Bacillus thuringiensis]	23	63	7.10E-29	322	296	g225559	1-333	Bt1G4369	1948 Bt1Gc5953	
[Bacillus subrilis] (L48554) vrrA gene product [Bacillus anthracis]	60	41	2.20E-20	241	213	g1162917	1-437	Bt1G4368	1947 Bt1Gc5937	
(Z99121) similar to ABC transporter (amino acid permease)	25	45	1.50E-27	310	292	g2635847	355-1	Bt1G4367	1946 Bt1Gc5949	
ORF IS231C [Bacillus thuringiensis]	25	50	1.10E-15	204	246	g225559	1-355	Bt1G4366	1945 Bt1Gc5948	
(U50744) BSMA [Bacillus stearothermophilus]	16	53	6.00E-15	199	269	g1255196	45-325	Bt1G4365	1944 Bt1Gc5942	
(ACONITASE) [Bacillus subtilis]									•	
ACONITATE HYDRATASE (CITRATE HYDRO-LYASE)	14	62	2.70E-33	373	314	g2506131	387-1	Bt1G4364	1943 Bt1Gc5939	
(AC006217) hypothetical protein [Arabidopsis thaliana]	27	75	3.30E-42	447	415	g4309751	520-191	Bt1G4363	1942 Bt1Gc5913	
I) [Bacillus caldolyticus] (AB013365) YlqF [Bacillus halodurans]	17	86	1.80E-18	223	223	g4514314	325-471	Bt1G4362	1941 Bt1Gc5918	
SIGNAL PEPTIDASE I (SPASE I) (LEADER PEPTIDASE	55	50	3.40E-24	277	266	g729934	1-301	Bt1G4361	1941 Bt1Gc5918	
SPORULATION PROTEIN N) [Bacillus subtilis]		ļ			• •		·			
ALANINE DEHYDROGENASE (STAGE V	26	54	2.90E-19	233	189	g585047	298-1	Bt1G4360	1940 Bt1Gc5917 Bt1G4360	
[Bacillus subtilis]										
(Z99119) similar to hypothetical proteins from B. subtilis	99	38	2.40E-07	118	115	g2635515	115-318	Bt1G4359	1939 Bt1Gc5903	
(AC007195) putative blue copper-binding protein II [Arabidopsis thaliana]) 8	ŏ	7.60E-27	302	203	g4>899/1	98-208	B[10433/	1938 Bt10c3899	
(ACOOTION) mutation blue common binding materia II		0 2	7 200 27	3 6	375	6460071	09.76			
(DS0AS3) von [Bacillus subtilis]	<u>.</u>	46	1 205-21	252	108	o1805463	377_1	B+1C4355	1037 R+1G-5880	
SYSTEM CARRIER PROTEIN (BRANCHED-CHAIN AMINO ACID UPTAKE CARRIER) [Bacillus subtilis]									٠.	
[] BRANCHED-CHAIN AMINO ACID TRANSPORT	27	63	6.70E-35	378	337	g3023410	1-395	Bt1G4354	1936 Bt1Gc5882	
nitrate reductase (EC 1.7.99.4) beta chain - Bacillus subtilis	10	63	3.60E-13	181	152	g2127147	1-151	Bt1G4353	1935 Bt1Gc5877	
(HEME SYNTHETASE) [Bacillus subtilis]				•			,			
FERROCHELATASE (PROTOHEME FERRO-LYASE)	. 4	28	6.60E-10	147	111	g417116	406-1	Bt1G4352	1934 Bt1Gc5841	
(HEME SYNTHETASE) [Bacillus subtilis]				į		ď				
FERROCHEL ATASE (PROTOHEME FERRO-LYASE)		ج هد	5 30E-19	228	190	9417116	1-368			
Synechocystis sp. [Synechocystis sp.] (AF015775) acetylornitine deacetylase 「Bacillus subtilis]	26	4	9.00E-17	· 213	238	g2415403	345-1	Bt1G4350	1932 Bt1Gc5838	
H+-transporting ATPase (EC 3.6.1.35) PMA1 -	∞	35	1.80E-05	113	. 71	g541317	475-1	Bt1G4349	1931 Bt1Gc5837	
ACETATE KINASE (ACETOKINASE) [Bacillus subtilis]	30	45	1.80E-16	209	213	g584719	1-365	Bt1G4348	1930 Bt1Gc5818	
HRCA [Bacillus stearothermophilus]									٠.	
NCBI gi description	Cyrg	Ident Cyrg	Prob	Score	nap Score	NCBI gi	Position	Gene Id	ID Contig Id	
	%	%	BlastP-	BlastP	aat_		:	:	SEQ	•
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	SEQ ID Contig Id Gene Id NO	Position	NCBI gi	nap Score	BlastP Score	BlastP- Prob	% % Ident Cvrg	%grv"g	NCBI gi description	
					•				PHOSPHOLYASE) (VEGETATIVE PROTEIN 216) (VEG216) [Bacillus subtilis]	
	1950 Bt1Gc5970 Bt1G4371	1-319	g1934614	110	159	2.70E-10	32	10	(U93874) cytochrome P450 102 [Bacillus subtilis]	
	Bt1Gc5971	1-267	g1813466	88	118	4.40E-07	34	38	(U61168) spore germination protein C2 [Bacillus firmus]	
	1952 Bt1Gc5979 Bt1G4373	1-291	g131720	95	121	5.00E-07	31	31	DIHYDROOROTATE DEHYDROGENASE (DIHYDROOROTATE OXIDASE) (DHODEHASE)	•
									[Bacillus subtilis]	
•	1953 Bt1Gc5992 Bt1G4374	1-358	g399058	216	264	8.00E-23	51	30	CHORISMATE SYNTHASE (5-	
					-				PHOSPHOLYASE) (VEGETATIVE PROTEIN 216)	
									(VEG216) [Bacillus subtilis]	
	1934 Bt1Gc3990 Bt1G43/3	1-004	g2492538	1/5	163	1.60E-11	3/	41	SULFAIE IRANSPURI AIF-BINDING PROTEIN CYSA [Synechocystis sn.]	
	1955 Bt1Gc5995 Bt1G4376	1-332	g3025120	137	172	5.50E-12	38	17	HYPOTHETICAL ABC TRANSPORTER ATP-BINDING	
	1956 Bt1Gc5998 Bt1G4377	1-385	g2226226	124	198	4.60E-15	34	27	PROTEIN YDIF [Bacillus subtilis] (Y14083) Hypothetical protein [Bacillus subtilis]	
	Bt1Gc6014	1-307	g1706723	126	154	2.60E-10	32	23	PUTATIVE EXODEOXYRIBONUCLEASE LARGE	-, -1
			٠.		à				SUBUNIT (EXONUCLEASE VII LARGE SUBUNIT) [Bacillus subtilis]	
	1958 Bt1Gc6030 Bt1G4379	19-441	g4512352	465	466	3.20E-44	60	28	(AB011836) proline transporter [Bacillus halodurans]	٠
	1959 Bt1Gc6035 Bt1G4380	477-66	g2829796	182	216	5.20E-17	33	29	CYTOCHROME D UBIQUINOL OXIDASE SUBUNIT I [Bacillus subtilis]	•
	1960 Bt1Gc6038 Bt1G4381	217-1	g2522016	150	184	7.00E-14	43	21	(AB007638) dehydrogenase [Bacillus subtilis]	
		246-1	g3183561	170	206	1.10E-16	52	31	VEGETATIVE PROTEIN 296 (VEG296) [Bacillus subtilis]	•
	Bt1Gc6046	264-1	g2618861	110	133	6.10E-09	38	51	(AF017113) putative acetyltransferase [Bacillus subtilis]	
	Bt1Gc6053	1-471	g2632105	584	607	3.60E-59	74	24	(Z98760) arginyl-tRNA synthetase [Arabidopsis thaliana]	
	Bt1Gc6055	450-1	g2577965	195	201	5.30E-15	34	20	(Y15254) PerA protein [Bacillus subtilis]	•
	ביים ביים ביים ביים ביים ביים ביים ביים	1-010	2007	. 10	. 60	1.100-13	۲	ą	ELEMENT IS232 [Insertion sequence IS232]	
	1966 Bt1Gc6071 Bt1G4387	192-17	g2984721	169	171	5.80E-13	2	91	(AF053927) probable spore germination protein D [Bacillus	
	1967 Bt1Gc6069 Bt1G4388	553-1	g1175719	44	469	1.50E-44	46	28	PENICILLIN-BINDING PROTEIN 3 (PBP 3) [Bacillus	
	1968 Bt1Gc6083 Bt1G4389	277-1	g586902	107	167	6.10E-12	3 8	26	HYPOTHETICAL 40.7 KD PROTEIN IN MECB-GLTX	
					% .	•	,		INTERGENIC REGION [Bacillus subtilis]	

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HYPOTHETICAL 50.0 KD PROTEIN IN SODA-COMGA INTERGENIC REGION [Bacillus subtilis]	36 22	3.10E-09	144	117	g1731036	54-350	Bt1G4436	8 Bt1Gc6405 Bt1G4436	2008
(Aro	. 60	1.90E-09	139	· 108	82413391	1-000	004400	/ PUTO0037/	7007
, (VE)		1 00E 00	3;	160	27415207	300 1	D+104434		3 1
_		5.80F-13	179	142	225559	1-214	Bt1G4434	6 Bt1Gc6401	2006
(Z75208) hypothetical protein [Bacillus subtilis]	45 . 9	0.034	%	109	g1770035	1-163	Bt1G4433	5 Bt1Gc6389	2005
_	40 54	2.10E-22	260	221	g1945657	423-1	Bt1G4432	4 Bt1Gc6386	2004
(AJ011676) DNA ligase [Bacillus stearothermophilus]	36 10	0.14	75	76	g3688229	1-207	Bt1G4431		2003
laminosum]							1))
(AJ224964) preprotein translocase [Phormidium	45 16	6.70E-10	152	168	g3288585	1-274	Bt1G4430	2002 Bt1Gc6373	200
	47 16	2.10E-28	328	312	g1934609	475-1	Bt1G4429	1 Bt1Gc6362	2001
cereus])
	40 25	3.50E-07	124	88	g3122814	1-283	Bt1G4428	0 Bt1Gc6334	2000
(AF05	56 · 98	0.013	48	139	g2984717	54-230	Bt1G4427	9 Bt1Gc6340	1999
	67 39	2.60E-49	514	562	g118716	1-442	Bt1G4426	8 Bt1Gc6333	1998
	99 93	1.90E-46	487	480	g540915	167-457	Bt1G4425	7 Bt1Gc6323	1997
	98 37	1.30E-06	111	233	g540914	1-161	Bt1G4424	7 Bt1Gc6323	1997
STAGE III SPORULATION PROTEIN E [Bacillus subtilis]	43 14	3.30E-17	222	185	g730776	326-1	Bt1G4422	6 Bt1Gc6304	1996
	54 36	3.40E-15	. 192	236	g2392192	426-178	Bt1G4421	5 Bt1Gc6299	1995
thermostable pullulanase [Bacillus stearothermophilus]	46 28	1.20E-44	470	431	g228654	558-1	Bt1G4420	4 Bt1Gc6294	1994
OXIDASE) []									
(COPROPORPHYRINOGENASE) (COPROGEN									
COPROPORPHYRINOGEN III OXIDASE									
PROBABLE OXYGEN-INDEPENDENT	41 27	2.30E-16	207	202	g1708181	300-1	Bt1G4419	3 Bt1Gc6286	1993
ORF IS231C [Bacillus thuringiensis]	65 9	1.60E-08	138	116	g225559	128-1	Bt1G4417	2 Bt1Gc6287	1992
(AL022603) VP1 like protein [Arabidopsis thaliana]	58 23	7.50E-34	376	372	g3080399	514-1	Bt1G4415	1 Bt1Gc6260	1991
(Y08031) cheD [Bacillus cereus]	57 27	3.60E-11	154	110	g1556378	24-183	Bt1G4414	0 Bt1Gc6277	1990
[Arabidopsis thaliana]		7.1						:	
(AC006217) putative retrotransposon polyprotein	75 10	6.20E-46	494	475	g4309763	382-1	Bt1G4413	9 Bt1Gc6275	1989
(AC006570) putative polyprotein [Arabidopsis thaliana]	88	4.50E-59	619	595	g4432807	1-401	Bt1G4412	8 Bt1Gc6274	1988
(gb D26601). [Arabidopsis thaliana]									
_	66 14	3.70E-28	322	159	g2342692	1-482	Bt1G4411	1987 Bt1Gc6265	198
retrotransposon-like gene (GB:AB00/467) [Arabidopsis thaliana]									•
									:
NCBI gi description	% % Ident Cvrg	BlastP- Prob I	BlastP Score	nap Score	NCBI gi	Position	Gene Id	Contig Id	8 5 8
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(Z99120) yunC [Bacillus subtilis]	89	40	3.80E-09	135	127	g2635733	271-1	4 Bt1G4459	2030 Bt1Gc6494
PROTEIN 2 IN GLVBC 3'REGION [Bacillus subtilis]						•			
HYPOTHETICAL ABC TRANSPORTER ATP-BINDING	18		3.40E-07	_	111	g1723607	327-1		
(7.97343) unnamed protein product [Arabidopsis thaliana]	25		0.023	. •	155	g2245111	1-266		
U DNA GYRASE SUBUNIT A Bacillus sp.]	. 14	35	4.20E-13	184	106	£4033397	1-354	8 Bt1G4455	2027 Bt1Gc6478
nitrate reductase (EC 1.7.99.4) beta chain - Bacillus subtilis	25	41	8.00E-19	233	187	g2127147	1-375	6 Bt1G4454	2026 Bt1Gc6466
subtilis]	, I.			•	· •		!		
(TRYPTOPHANTRNA LIGASE) (TRPRS) [Bacillus	•			٠					
TRYPTOPHANYL-TRNA SYNTHETASE	22	45	1.30E-10	154	121	g135188	47-267	5 Bt1G4453	2025 Bt1Gc6465
(AF147259) No definition line found [Arabidopsis thaliana]	18	63	5.30E-10	146	101	g4773910	1-218	7 Bt1G4452	2024 Bt1Gc6457
(Y11170) BC541A protein [Bacillus cereus]	45	45	2.00E-12	166	130	g2462088	162-412	0 Bt1G4451	2023 Bt1Gc6460
subtilis]									
(Z82044) unidentified transporter-ATP binding [Bacillus	20	35	6.90E-07	124	136	g1673392	1-347	6 Bt1G4450	2022 Bt1Gc6456
(Y14078) Hypothetical protein [Bacillus subtilis]	42	39	2.90E-18	221	141	g2226127	395-1	5 Bt1G4449	2021 Bt1Gc6435
(AL035539) putative protein [Arabidopsis thaliana]	11	54	4.00E-11	163	87	g4539351	138-422	2 Bt1G4448	2020 Bt1Gc6452
(AC003981) F22O13.10 [Arabidopsis thaliana]	15	89	6.20E-83	834	796	g3063448	556-1	9 Bt1G4447	2019 Bt1Gc6449
[Bacillus subtilis]									
DEHYDROGENASE (COLD SHOCK PROTEIN CSI14)									
2,3-DIHYDRO-2,3-DIHYDROXYBENZOATE	52	68	1.90E-39	421	445	g1169301	1411	7 Bt1G4446	2018 Bt1Gc6447
from this gene. [Arabidopsis thaliana]									
adaptin 1 from Arabidopsis thaliana. EST gb H37393 comes									
(AC005966) Strong similarity to gb AF061286 gamma-	ω	83	4.30E-07	128	92	g4249386	1-463	5 Bt1G4445	2017 Bt1Gc6445
(AC003000) hypothetical protein [Arabidopsis thaliana]	15	90	1.50E-51	540	548	g2642163	74-436	3 Bt1G4444	2016 Bt1Gc6443
Bacillus subtilis []			٠						-
aspartate-semialdehyde dehydrogenase (EC 1.2.1.11) -	. 40	32	6.80E-15	193	148	g538921	1-433	7 Bt1G4443	2015 Bt1Gc6437
[Arabidopsis thaliana]								···•	
(AC006250) putative Athila retroelement ORF1 protein	7	8	1.60E-26	310	305	g4263543	1-201	3 Bt1G4442	2014 Bt1Gc6433
(X02441) gtg start [Nicotiana tabacum]	98	· 86	2.40E-14	184	167	g1334366	219-92	8 Bt1G4441	2013 Bt1Gc6428
longistaminata				•					•
(U72725) receptor kinase-like protein [Oryza	26	44	1.10E-22	271	223	g2586081	86-532	1 Bt1G4440	2012 Bt1Gc6421
NADH dehydrogenase 2-like ORF 180 [Nicotiana tabacum]	29	65	1.50E-05	102	121	g225268	471-310	0 Bt1G4439	2011 Bt1Gc6420
nitrate reductase delta chain - Bacillus subtilis []	59	30	.7.00E-08	123	109	g1075873	362-37	9 Bt1G4438	·2010 Bt1Gc6419
				-					
purine nucleoside phosphorylase (EC 2.7) II - Bacillus	66	45	4.90E-32	351	338	g2126871	463-1	5 Bt1G4437	2009 Bt1Gc6415
	ď				Score				NO
NCBI gi description	\cap	∓	Prob	Score	nap	NCBI gi	Position	I Gene Id	ID Contig Id
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Table 1

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Bt1G4492	Bt1G4491	Bt1G4486	Bt1G4485		Bt1G4484	00110110	<u>.</u>	Bt1G4482		Bt1G4481		Bt1G4480	Bt1G4478	Bt1G4476	Bt1G4474		Bt1G4473		Bt1G4472	Bt1G4471		Bt1G4469		Bt1G4468	Bt1G4467	Bt1G4466		Bt1G4465	Bt1G4464		Bt1G4461	Bt1G4460	сепе та	}
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g458	g228	g229	g229	•	2249	8400	27.5	g321919		g194		g229	g483	g231	g170		g136144		g467	g252		g131137		ğ173	g456	g261	C	E354	g134761		g730965	g467	. 2	Ś
g4584140	g2286111	g2293212	g2293215		£2497382	84220241	3	919		g1944414	•	g2293148	g4835229	g2314492	g1703404		144		g4678307	g2529467	,	137		g1731026	g4567262	g2618863		g3548811	761		965	g4678942	NCBI 81	Ē 1
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130	130	254	112		119		3	83		242		242	470	101	504		266		256	256		240		148	353	256		294	341		214	230	nap Score	_,RR
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145	196	299	112		154	110	100	106		257		248	427	122	503		307		345	326		321		147	336	320		320	383		211	293	ore	stP
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7.10E-10	1.30E-15	1.60E-26	1.70E-05		2.40E-10	2.50E-00	305	4.50E-06		6.00E-21		4.00E-21	4.30E-40	8.00E-07	3.80E-48		4.10E-27		4.20E-31	2.20E-29		7.20E-28		2.60E-09	1.90E-30	9.40E-29		9.40E-29	2.00E-35		4.00E-16	6.80E-26	Prob	BlastP-
io	<u>:</u> 5	26	S		Ö	Ġ	₹ .	9		21		21	6	97	48		27		<u>3</u>	29		28		9	<u>3</u> 0	29	ļ	29	35		6	26	-	•
38	55	47	29		35	26	3	39		4		52	54	30	49		66		87	72		48		35	ထ	45	:- !	2	48		53	61	Ident	%
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(AJ010139) fumarate hydratase [Bacillus cereus]	(U7889	(AF008)	(AF008220) YtpT [Bacillus subtilis]	ELEMENT IS232 [Insertion sequence IS232]	[Staphylococcus aureus] TRANSPOSASE FOR INSERTION SEOUENCE		plasmid	hypothetical 16.9K protein - Salmonella typhimurium	stearothermophilus]	(D87026) glycogen phosphorylase [Bacillus	subtilis]	(AF008220) dihydroxynaphthoate synthase [Bacillus	(AL049862) putative protein [Arabidopsis thaliana]	AE0	Ħ	ELEMENT IS231C [Bacillus thuringiensis]	TRANSPOSASE FOR INSERTION SEQUENCE	[Arabidopsis thaliana]	(AL049655) vacuolar protein sorting-like protein	(AF006665) YokS [Bacillus subtilis]	APOPROTEIN A1 [Pisum sativum]	PHOTOSYSTEM I P700 CHLOROPHYLL A	INTERGENIC REGION [Bacillus subtilis]	3	(AC006841) putative ubiquitin [Arabidopsis thaliana]	(AF017	thaliana]	(AC005)	STAGE II SPORULATION PROT	(RELAXING ENZYME) (UNTWISTING ENZYME)	DNA TOPOISOMERASE I (OMEGA-PROTEIN)	(AL049)		
0139	39)822)822		SPC S		id N	hetic	then	26)	<u>s</u>)822	1986	2063	MA	EZ	SPC	idop	1965)666	ŘOJ	SOS	RGE	HE	2684		a)531	E		TOP	1971		
) fu	 MADS box protein [Oryza sativa] 	220) YtpQ [Bacillus subtilis]	0) Y		SAS(1	NTP16 []	al 16	noph	glyco) Ei	2) p	4) fu	TE 5	r IS2	SAS	sis th	5) va	5) Y	EZ	STE	ZIC	II.	בי פר	[13] YvpB [Bacillus subtilis]	,	3) ki	SPO	GE	OIS	1) SE		
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						(Ar 101234) D-alaillie-D-alailyi carrier protein ilgase DitA		E E				llus	نق	ri 26	SHIKIMATE 5-DEHYDROGENASE [Bacillus subtilis]									Ċ.	ana]			rabi	ns si	Ð		thali		
						7) * *	•						(AE000634) fumarase (fumC) [Helicobacter pylori 26695]	is]									HYPOTHETICAL 73.2 KD PROTEIN IN SODA-COMGA			1	313) kinetochore (SKP1p)-like protein [Arabidopsis	II SPORULATION PROTEIN D [Bacillus subtilis]			711) 5B protein like protein [Arabidopsis thaliana]	-	
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Table 1

2076 Bt1Gc6876 2077 Bt1Gc6881		2074 Bt1Gc6872	2073 Bt1Gc6871	2072 Bt1Gc6849	2071 Bt1Gc6842		2070 Bt1Gc6837	2069 Bt1Gc6819	2068 Bt1Gc6836		2067 Bt1Gc6801	2066 Bt1Gc6802		2065 Bt1Gc6798		2064 Bt1Gc6778		2063 Bt1Gc6786		2062 Bt1Gc6777		2061 Bt1Gc6754	2060 Bt1Gc6729	2059 Bt1Gc6724		2058 Bt1Gc6709	2057 Bt1Gc6703	2056 Bt1Gc6700	2055 Bt1Gc6686		NO County or
Bt1G4525 Bt1G4526	_	Bt1G4522	Bt1G4521	Bt1G4518			Bt1G4515	Bt1G4514	Bt1G4513		Bt1G4508	Bt1G4507		Bt1G4506		Bt1G4505		Bt1G4504		Bt1G4503		Bt1G4502	Bt1G4501	Bt1G4500		Bt1G4499	Bt1G4496	Bt1G4495	Bt1G4494		Conc.
1-564 1-641	1-629	1-605	129-405	508-403	1-630		1-342	1-392	207-1		525-1	214-1		406-1		161-373		1-169		1-435		292-1	287-1	374-1		1-275	1-592	120-1	551-1	1-347	ı oğudu
g4583548 g4415926	g3522943	g3646371	g1002380	g2529340	g4914343		g3123297	g1840129	g3915204		g3915461	g2293210		g1731052	1	g4835325	٠	g3328184		g2811062		g482286	g2635764	g2443256		g729583	g2130141	g97193	g479357	g2832244	. 6
655 330	557	163	93	174	310		130	148	95		272	135		89		102		139		126		169	103	205		119	254	96	235	367	Score
645 377	592	208	56	190	384		118	180	111		308	141		142		109		148		178		204	145	221		121	184	105	251	183	Score
3.40E-63 1.40E-33	2.00E-56	6.90E-17	_	5.60E-15	2.70E-34		4.90E-09	8.10E-14	3.20E-05		8.50E-27	4.30E-09		9.00E-09		9.40E-06		4.00E-10		7.60E-13		6.40E-15	3.30E-10	1.90E-17		6.10E-07	4.20E-13	9.60E-05	1.60E-20	2.00E-13	Prob
4 3 73	57	53	38	97	43		37	37	48		37	43	_	32		3 8		59		35		4	39	39		35	51	51	65	63	Ident (
51 20	16	60	26	14	19		- 18	45	7		30	20		17		24		19		29		7	63	24		27	14	6	14	25	Cvrg
(AJ010820) FtsY homolog [Arabidopsis thaliana] (AC006282) unknown protein [Arabidopsis thaliana]	(AC004411) putative p-glycoprotein [Arabidopsis thaliana]	elegans] (AJ004916) chorismate mutase precursor [Prunus avium]	(U24189) RRM-type RNA binding protein [Caenorhabditis	(L81162) homologue; putative [Zea mays]	(AC005489) F14N23.29 [Arabidopsis thaliana]	[Bacillus subtilis]	EXCINUCLEASE ABC SUBUNIT B (DINA PROTEIN)	(U86347) beta-lactamase [Cloning vector pCALnFLAG]	EXCINUCLEASE ABC SUBUNIT A [Bacillus subtilis]	INTERGENIC REGION [Bacillus subtilis]	HYPOTHETICAL 66.6 KD PROTEIN IN PURD-SAPB	(AF008220) YtoP [Bacillus subtilis]	SYNTHASE (DXP SYNTHASE) [Bacillus subtilis]	PROBABLE 1-DEOXYXYLULOSE-5-PHOSPHATE	[Streptomyces coelicolor]	(AL049863) putative transcriptional regulator	pQE30NST]	(AF074376) beta lactamase precursor [Cloning vector	GLYCOGEN) SYNTHASE) [Bacillus stearothermophilus]	GLYCOGEN SYNTHASE (STARCH (BACTERIAL	(version 2) - Bacillus subtilis []	subtilis] DNA-directed DNA polymerase (EC 2.7.7.7) III alpha chain	(Z99120) similar to NifU protein homolog [Bacillus	(D86417) YfmM [Bacillus subtilis]	subtilis]	GLYCOGEN BIOSYNTHESIS PROTEIN GLGD [Bacillus	mudrA protein - maize transposon MuDR [Zea mays]	maysj leukotoxin B - Pasteurella haemolytica []	hypothetical protein 612 - maize transposon MuA2 [Zea	(AF031569) hypothetical protein [Zea mays]	Tropa G. acost peron

8 pheromone cAD1 binding protein precursor - Enterococcus	4	4.80E-14	190	147	g1075694	1-311	2100 Bt1Gc7041 Bt1G4576	210
	37 21	1.50E-06	. 118	105	g3290176	225-1	99 Bt1Gc7037 Bt1G4575	2099
•	41 11	6.00E-05	106	52	g2635778	267-1	98 Bt1Gc7023 Bt1G4574	2098
	64 31	1.00E-47	499	515	g1352830	1-522	97 Bt1Gc7015 Bt1G4573	2097
PREPROTEIN TRANSLOCASE SECA SUBUNIT	30 _, 16	3.80E-06	119	128	g134396	1-401	96 Bt1Gc7002 Bt1G4570	2096
	86 70	4.80E-80	804	738	g2529340	1-536	95 Bt1Gc6994 Bt1G4568	2095
4 ORF IS231C [Bacillus thuringiensis]	71 24	5.80E-36	388	372	g225559	19-362	94 Bt1Gc6974 Bt1G4567	2094
5 STAGE V SPORULATION PROTEIN T [Bacillus subtilis]	44 35	7.80E-09	132	106	g586883	1-188	93 Bt1Gc6980 Bt1G4565	2093
roseus 7 ORF IS231C [Bacillus thuringiensis]	71	1.80E-06	119	104	g225559	100-1	92 Bt1Gc6965 Bt1G4564	2092
	52 40	7.70E-50	519	462	g3954807	1-625	91 Bt1Gc6957 Bt1G4562	2091
_					•			
formyltransferase [Bacillus cereus] (Z81356) UDP-N-acetylglucosamine 1-	48 29	2.60E-26	297	295	g1648861	377-1	90 Bt1Gc6951 Bt1G4561	2090
	59 30	1.10E-25	291	246	g2072373	1-308	39 Bt1Gc6956 Bt1G4558	2089
5 (AL021960) putative protein [Arabidopsis thaliana]	38 . 45	8.00E-31	254	184	g2911073	251-652	38 Bt1Gc6925 Bt1G4553	2088
8 (AL035526) extensin-like protein [Arabidopsis thaliana]	49	0.00085	97	57	g4539386	275-474	37 Bt1Gc6929 Bt1G4551	2087
_	50 54	4.20E-26	295	289	g4584429	198-612	36 Bt1Gc6923 Bt1G4550	2086
thaliana]								
(version 2) - Bactilus submits [] (AC005967) putative carbonyl reductase [Arabidopsis	40 53	5.30E-26	294	201	g4115379	142-630	35 Bt1Gc6927 Bt1G4549	2085
/ UNA-directed UNA polymerase (EC 2.7.7.7) III alpha chain	48	1.10E-15	211	141	g482286	1-087	54 BITCC0922 BITC4548	2084
	66 47	2.80E-75	759	719	g4056432	1-619	Bt1Gc6912	2083
_		2.10E-15	194	148	g3341694	280-571	Bt1Gc6910	2082
_	29 14	4.30E-08	138	101	g2959781	1-409	31 Bt1Gc6909 Bt1G4544	2081
INTERGENIC REGION [Bacillus subtilis]								
[Alabidopsis manana] 2 HYPOTHETICAL 22.6 KD PROTEIN IN EPR-GALK	39 52	1.00E-13	178	149	g732345	1-310	2080 Bt1Gc6902 Bt1G4542	208
	39 15	4.70E-07	127	113	g4454463	1-652	79 Bt1Gc6891 Bt1G4539	2079
3 (AC006068) putative ch-TOG protein [Arabidopsis	6	0.00031	105	70	g4263790	529-367	78 Bt1Gc6890 Bt1G4537	2078
	TOUT CALE	1100	2000	æ				NO
NCBI gi description		1-	BlastP	aat_	NCBI gi	Position	Q Contig Id Gene Id	SEQ

Table 1

2121 2121 2122	2119		2118	2117	2116		2115		21,14	2113	2112	2111		2110		2109		2108	1	2107	2106	3105	2104	. 1	2103	3103	2101	C	SEQ
Bt1Gc7273 Bt1Gc7277	Bt1Gc7236		Bt1Gc7235	Bt1Gc7224	Bt1Gc7187		Bt1Gc7182		Bt1Gc7175	Bt1Gc7174	Bt1Gc7164	Bt1Gc7137		Bt1Gc7125		Bt1Gc7121		Bt1Gc7113	9	B+10-7110	Bt1Gc7103	B+1.G-7077	Bt1Gc7072		Br1Gc7064	B+10-7056	Bt1Gc7050		Contig Id
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Bt10	Bt1C		Bt1C	Bt1G	Bt1G		Bt1G		Bt1G	Bt1G	Bt1G	Bt1G		Bt1G		Bt1G		Bt1G	5	ביות הוא			Bt1G		בי בי בי		Bt1G		Gen
Bt1G4598 Bt1G4598	Bt1G4596		Bt1G4595	Bt1G4594	Bt1G4593		Bt1G4592		Bt1G4591	Bt1G4590	Bt1G4589	Bt1G4588	•	Bt1G4587		Bt1G4586		Bt1G4585	1	R+1G4584	B+1G4593	B+1 C4 58 1	Bt1G4580	1000	B+1G4570	B+1 C4578	Bt1G4577	:	Gene Id
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343-1 301-1 1-420	1-430		360-70	1-125	1-416		166-1		349-1	546-1	385-1	1-183		1-387		1-313		1-278	1.1	1-171	46.484	163_1	547-1	1-67	1506	1-340	253-1		Position
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g2293126 g135811 g1934652	g2632017	•	g120717	g2493761	g2635199		g3915700		g2984720	g556177	g1304006	g2274867		g1176995		g4467246		g141397	8117700	6113368	5370	~1651078	g732409	B4700701	0230	73070573	g126772		NCBI gi
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7.10E-08 6.90E-10	1.30E-22		1.70E-20	1.60E-08	2.50E-10		7.30E-06		2.60E-24	5.50E-56	4.20E-11	2.00E-13		.30E-22		2.60E-09		2.10E-09	Č	0.00E-05		7 00E-11	1.70E-45	Į.	1 305-07	1 705 07	1.20E-08		BlastP- Prob
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(U89962) beta-lactamase [Cloning vector pVP16] THREONINE SYNTHASE [Bacillus sp.] (U93876) amino acid transporter [Bacillus subtilis	stearothermophilus] (AJ002571) DppE [Bacillus subtilis]	(PHOSPHOGLUCOSE ISOMERASE A) [Bacillus) DÚ	NOT	(Z99117) similar to	DEHYDROGENASE (GAPDH) [Bacillus	GLYCERALDEHYDE 3-PHOSPHATE	cereus]	(AF053	(M2262)	(D84432) SpoVAF [Bacillus subtilis]	(AJ000005) glucose kinase [Bacillus megaterium]	PROTEIN IN IDH 3'REGION [Bacillus subtilis]	P01	coelicolor]	(AL035)	INTERGENIC REGION (ORF238) [Bacillus subtilis]	HYPOTHETICAL OXIDOREDUCTASE IN RTP-PELB	ביול ביונים	AT COHOL DEHADS OCENIAGE LADA D. 12	(D)000(5) linearstein NIAD [Cymechocyctic en]	ORFK)	HYPOTHETICAL 36.3 KD LIPOPR		ATP_DEPENDENT OF PROTE ACE ATP_BINDING	crassa]	MALA'	-	
6) at 12	S71)	РНО)SE-	g.	7) sir		RA			?1) is	2) Sp	05)	Ï	HE	9	569)	GEN		֡֟֝֟֓֓֓֓֓֓֓֟֟֓֓֓֓֓֟֟֓֓֓֓֟֟֓֓֓֓֓֓֟֓֓֓֟֟֓֓֓֟֓֓֓֓	֓֞֝֞֜֜֞֜֜֜֝֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓	クとぎき	֝֟֟֝֟֝֟֝֟֝֟֝֟֝ <u>֚֚֚֚֚֚</u>	HE		(בוקה) (ביין	3	piasi TE S	1	
beta-lactamase [Cloning vector pVP16]NINE SYNTHASE [Bacillus sp.]amino acid transporter [Bacillus subtilis]	Dpp.	GLU	6-PE	Ď	nilar		DE		prot	1) isocitrate lyase [Escherichia coli]	7V0C	gluc	Z U	TICA		pute	IC R	TICA	Į.	יות קייני		[Bacillus subtilis]	TICA		PENDENT CI P PROTEASE ATP-B		Plasmid PAUT Plasmid PAUT TE SYNTHASE, GLYOXYSON		٠
YNT acid		ĊŎ	ISO	OHS	40 P	ASE	T T		oable	ate l	=	ose k	H 3"	LA		tive	EGI	D J			4010	ıs su	T 36	7 7	Z	- 5 -	YHY.		Z
hase HAS tran	Bacil	SE IS	AHG	S	ypot	(c))E 3-		spo	yase	Baci	cinas	REG	BC		pho	8	Ä	Š			btilis	5.3 K	֝֞֞֜֜֜֝֜֝֜֝֜֜֝֝֓֓֓֜֝֜֜֜֝֓֓֓֓֓֓֓֓֓֓֓֓֓֓֜֝֓֓֓֡֝֡֓֜֝֓֡֓֓֡֝֡֡֡֝֡	ם זיר קינו	1	SE, C	<u>.</u> 3	СВІ
spor	lus s	Š	ΙΞΙ	PRO	hetic	APU USA	Ħ		re ge	[Es	llus	ë H	Ö	TRA		spho	(OR	ORE	7		7 E		D S		ַלְבָּלְינָ הַלְבָּלְינָ	3	ATC		6
onung Baci ter [ubtil	ER	SON	TEI	al p	_ H	SPF		E E	cher	subti	3acil	Œ	NSP		diest	F238	Ü	70	A CIE	3	Ģ	IPO	֓֞֞֜֜֞֜֜֞֜֜֜֓֓֓֓֜֜֜֜֜֓֓֓֓֓֜֜֜֜֜֓֓֓֓֓֜֜֜֜֓֓֓֡֓֜֡֓֜			OX P		escr
llus :	is]	SE	ER	Ω CS	rotei	Baci	IAT		natio	ichia	lis]	lus n	zcill	OR1		eras	ш Ш	CTA	3				PRO	֓֞֞֜֜֜֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֡֓֓֓֓֓֡֓֜֓֡֓֡֓֡֓֡֓֡֓֡֓֡֓֡֡֡֡֡	N S	5	SO	•	NCBI gi description
Sp.]	•	<u>8</u>	ASE	PA	ns []	llus	; ITI		n pro	coli		negai	ns su	ER		e S	acill	SE I		DE S			TEN	1	A To		MAI –	_	5
subt		Baci	∑ . ∀	Вас	Bacil				otein	_		teriu	btilis	ATP		шерt	ns su	Z	7	7 G	3	.	V PR	Į.	֓֞֞֞֓֞֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֡֓֓֓֓֓֡֓֓֡	<u>.</u>	Z		
)	llus	}PI /	illus	lus s				C			3	ت	BIP		569) putative phosphodiesterase [Streptomyces	ıbtili	TP-P	ýmc		3	_	ECU		7		euro		
		,	GLUCOSE-6-PHOSPHATE ISOMERASE A (GPI A)	cere	(Z99117) similar to hypothetical proteins [Bacillus subtilis]				927) probable spore germination protein C [Bacillus					HYPOTHETICAL ABC TRANSPORTER ATP-BINDING		ces	ည	ELB	TI CIL	AT COHOL DEHADBOCENIACE I (ADH I) (Zymomomos			HETICAL 36.3 KD LIPOPROTEIN PRECURSOR	2	ถื		MALATE SYNTHASE, GLYOXYSOMAL [Neurospora		•
				us]	<u>s</u>				lus				• •:	G				•.	8	9			Ħ				-		
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Table 1

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2142 Bt1Gc7426 Bt1G4619	2141 Bt1Gc7400 Bt1G4618	2140 Bt1Gc7405 Bt1G4617		2139 Bt1Gc7396 Bt1G4616	2138 Bt1Gc7397 Bt1G4615	.*	2137 Bt1Gc7374 Bt1G4614	ζ		2136 Bt1Gc7375 Bt1G4613	2135 Bt1Gc7361 Bt1G4612	2134 Bt1Gc7357 Bt1G4611		2133 Bt1Gc7352 Bt1G4610		2132 Bt1Gc7347 Bt1G4609	2131 Bt1Gc7345 Bt1G4608	•	2130 Bt1Gc7350 Bt1G4607	2129 Bt1Gc7337 Bt1G4606		2128 Bt1Gc7324 Bt1G4605	2127 Bt1Gc7325 Bt1G4604	2126 Bt1Gc7322 Bt1G4603		2125 Bt1Gc7312 Bt1G4602	2124 Bt1Gc7286 Bt1G4601	2123 Bt1Gc7295 Bt1G4600	ID Contig Id Gene Id	
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403-1	530-1	334-1		1-308	248-48		218-1	:	: . *	345-1	1-351	1-489		438-1		1-375	1-430		1-174	1-236		229-1	315-1	214-391		203-1	1-370	1-356	Position	•
g2492639	g2415745	g2635763		g482286	g3955202		g2492562			g1346442	g2497382	g1352095		g482286	1	g1175720	g2149596	,	g2497382	g4584142		g114646	g114636	g114510	•	g114646	g4584142	g2618835	NCBI gi	
140	211	227		186	115		140			95	174	212		263		152	8 7		130	169:		119	122	110	•	89	319	203	nap Score	aat_
) 210	281	7 265		5 220	· . 69) 116	•		130	1 229	2 174		3 285			7 139) 155): 145	-	128	! 132	174		110	349	166		BlastP
2.50E-16	1.30E-24	2.00E-22		1.30E-16	0.081		3.90E-07			1.40E-07	1.50E-18	1.00E-12		1.50E-23		3.00E-16	1.90E-08		1.90E-10	8.90E-10		7.00E-08	2.50E-08	2.20E-12		6.80E-06	7.90E-32	4.10E-12	Prob	BlastP-
υ ω	40	2 53		6 46	39		7 38			7 29	49	38		4		_	\$ 47		56	47		40	37	 2		38	2 57	. 42	Ħ	\$
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3-ISOPROPYLMALATE DEHYDRATASE LARGE SUBUNIT (ISOPROPYLMALATE ISOMERASE) (ALPHA-IPM ISOMERASE) (IPMI) [Actinoplanes teichomyceticus]	(AB000617) YeeH [Bacillus subtilis]	(Z99120) similar to hypothetical proteins [Bacillus subtilis]	(version 2) - Bacillus subtilis []	DNA-directed DNA polymerase (EC 2.7.7.7) III alpha chain	(AF022796) MoaB [Staphylococcus carnosus]	mutans]	EXCINUCLEASE ABC SUBUNIT A [Streptococcus	SYNTHETASE) [Anabaena PCC7120]	ISOPROPYLMALATE SYNTHASE) (ALPHA-IPM	ELEMENT IS232 [Insertion sequence IS232] 2-ISOPROPYLMALATE SYNTHASE (ALPHA-	ICE	COMG OPERON PROTEIN 1 [Bacillus subtilis]	(version 2) - Bacillus subtilis []	DNA-directed DNA polymerase (EC 2.7.7.7) III alpha chain		HYPOTHETICAL OXIDOREDUCTASE IN PBPC-LRPC	(U89796) chitinase [Bacillus thuringiensis]	ELEMENT IS232 [Insertion sequence IS232]	cereus] TRANSPOSASE FOR INSERTION SEQUENCE	(AJ010139) DNA alkylation repair enzyme [Bacillus	[thermophilic bacterium PS3]	ATP SYNTHASE GAMMA CHAIN PRECURSOR	ATP SYNTHASE GAMMA CHAIN [Bacillus megaterium]	ATP SYNTHASE ALPHA CHAIN [Bacillus megaterium]	[thermonhilic hacterium DC2]	cereus] ATP SYNTHASE GAMMA CHAIN PRECURSOR	(AJ010139) DNA alkylation repair enzyme [Bacillus	(AF017113) cell division protein [Bacillus subtilis]	NCBI gi description	

Table 1

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2163 Bt1Gc7654 Bt1G4641		2162 Bt1Gç7653 E	2161 Bt1Gc7651 E	2160 Bt1Gc7625 B		2159 Bt1Gc7619 F	2158 Bt1Gc7616 B	2157 Bt1Gc7597 B	2156 Bt1Gc7590 E			Bt1Gc7592	2154 Bt1Gc7583 B		2153 Bt1Gc7507 B		2152 Bt1Gc7508 B	•	Bt1Gc7500	Bt1Gc7495	2150 Bt1Gc7495 B		Bt1Gc7498	2148 Bt1Gc7488 B		Bt1Gc7473	2146 Bt1Gc7470 B	2145 Bt1Gc7392 B	2144 Bt1Gc7440 B	2143 Bt1Gc7429 B	SEQ ID Contig Id O
3t1G4641		Bt1G4640	Bt1G4639	Bt1G4638		Br1G4637	Bt1G4636	Bt1G4635	Bt1G4634	-		Bt1G4633	Bt1G4632		Bt1G4631		Bt1G4630		Bt1G4629	Bt1G4628	Bt1G4627		Br1G4626	Bt1G4625		Rt1G4624	Bt1G4623	Bt1G4622	Bt1G4621	Bt1G4620	Gene Id
492-1		1-186	416-1	254-1		255-1	363-1	62-274	421-1			265-1	1-311		168-1		1-223		1-540	317-504	1-275	,	1-533	1-547		1-268	143-1	1-264	1-423	1-272	Position
g584920 ,		g3183527	g2633126	g2635763	O C	9585047	g3122850	g2636123	g3914289			g728788	g2815006	•	g2497382	(g2501412	9	g3925779	g2119971	g1945118	0.000	g1351995	e549706	0.00	o135106	g1934831	g2293164	g1731004	g72992	NCBI gi
723		104 104	168	156	,	<u>1</u> 06	184	128	216			177	192		130		124		238	247	373		20 2	120	į	<u> </u>	100	144	262	179	nap Score
493		120	224	177	į	148	204	188	197			208	230	,	187		155		181	276	372	į	ر کر	205		140	119	191	297	131	BlastP Score
4.40E-47		1.50E-07	6.80E-18	9.00E-13	9. 10	8 40F-10	1.10E-15	9.10E-15	1.20E-14			6.00E-16	5.10E-18		6.20E-14		1.70E-10		5.00E-13	2.20E-23	2.90E-34		3 70F-49	1.50E-16):00E	9 80F-09	9.20E-07	1.90E-14	2.60E-26	5.10E-08	BlastP- Prob
89		40	36	40	Ç	بر 0	40	50	4	٠		50	41	٠.,	<u>61</u>		47	ļ	29	8	83	ć	<u>ب</u>	35	į	4	52	53	46	49	% Ident (
30		39	30 (18		<u>ي</u>	26]	77 (21 '		_	15 ,	13 (13				31 (12 1	%		_	45		_		22 (44	26 1	Cvrg
60 KD CHAPERONIN (PROTEIN CPN60) (GROEL	(TRANSCRIPT CLEAVAGE FACTOR GREA) (GENERAL STRESS PROTEIN 20M) (GSP20M) [Bacillus	subtilis] TRANSCRIPTION ELONGATION FACTOR GREA	(Z99108) similar to RNA methyltransferase [Bacillus	(Z99120) similar to hypothetical proteins [Bacillus subtilis]	SPORULATION PROTEIN N) [Bacillus subtilis]	Bacillus firmus ALANINE DEHYDROGENASE (STAGE V	PREPROTEIN TRANSLOCASE SECA SUBUNIT	(Z99122) ywsA [Bacillus subtilis]	TOPOISOMERASE IV SUBUNIT B [Bacillus subtilis]	(ACETYL-COA SYNTHASE) [Bacillus subtilis]	COA LIGASE) (ACYL-ACTIVATING ENZYME)	ACETYL-COENZYME A SYNTHETASE (ACETATE	(AJ001087) pullulanase [Thermotoga maritima]	ELEMENT IS232 [Insertion sequence IS232]	TRANSPOSASE FOR INSERTION SEQUENCE	[Methanococcus jannaschii]	TRYPTOPHAN SYNTHASE BETA CHAIN	resistance protein [Schizosascharomyses nombe]	(AL034353) putative major facilitator family multi-drug	molecular chaperone 60 GroEL - Bacillus sp []	(D88802) groES [Bacillus subtilis]	[Schizosaccharomyces nombe]	PROBABLE CALCILIM-TRANSPORTING ATPASE	KTI12 PROTEIN (Saccharomyces cerevisiae)	TOVER TO VELL CONTROL DESIGNATION (OF CONTROL DE	GI ITAMVI -TRNA SVNTHETASE (GI ITAMATE-	(Z93939) unknown [Bacillus subtilis]	(AF008220) SAM synthase [Bacillus subtilis]	LYTB PROTEIN HOMOLOG [Bacillus subtilis]	recE protein - Bacillus subtilis []	NCBI gi description

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	(U24659) JadR1 [Streptomyces venezuelae] (AB011836) similar to B.subtilis ywgB gene(27%-identity) [Bacillus halodurans]	. 49) 40 7 42	6.40E-10 3.90E-07	143 116	80 102	g886038 g4512355	381-82 198-1	Bt1G4663 Bt1G4664	3 Bt1Gc7804 4 Bt1Gc7809	2183 2184
	GTP-BINDING PROTEIN TYPA/BIPA HOMOLOG	21	63	4.30E-40	427	400	g3183185	1-384	Bt1G4662	2 Bt1Gc7791	2182
	(D78193) yydK [Bacillus subtilis]	45	28	7.50E-06	108	101	g1064785	1-327	Bt1G4661		2181
	(AJ005075) enzyme I [Bacillus megaterium]	∞	68	2.10E-08	138	116	g3021327	1-131	Bt1G4660	Bt1Gc7778	2180
	(299107) similar to acritiavin resistance protein [bacilius	11	00	9.005-03	10/	70	82032763	040-1	0104039	bring.	2117
	(J200107) : iactate denydrogenase [Bacillus subtilis]	: 2		0.90E-07	103	211	g18033//	1 2/0	Bt1C4658		2170
	FACTOR NIRA [Emericella nidulans]	3	•		1	:	21005277	1	D+10/4650		2
	NITROGEN ASSIMILATION TRANSCRIPTION	∞	35	6.40E-11	164	94	g128340	1-561	Bt1G4657	7 Bt1Gc7760	2177
	(U51115) unknown protein [Bacillus subtilis]	14	42	1.10E-08	139	127	g2239289	188-1	Bt1G4655	5 Bt1Gc7743	2176
	[Bacillus subtilis]										
	PROBABLE PHOSPHOMANNOMUTASE (PMM)	15	36	2.40E-07	128	115	g2851553	259-1	Bt1G4654	5 Bt1Gc7719	2175
	Plegans										
×	R09H10.3 IN CHROMOSOME IV [Caenorhabditis	2	۲	2.305-03	130	72	80710174	717-21	נייטרטיט		. 1
· ·	LANDOTHETTO AT TO A NICTUAL PROTECTION OF THE PR	21	-	3 905 00	136	3 6	8123733 2015104	542-271	B+1G4652		2174
	I ACTOOR DEDMEACE [Kling/aromy/cas lactic])	် ၁	1 305-15	205	168	9125035	471_1	B+1C4653	R+1G-7717	2173
	[Bacillus halodurans]			!	;	1	G				
	(AB017508) fus homologue (identity of 87% to B. subtilis)	6		1.70E-14	196	196	g4512401	257-376	Bt1G4651		2172
	(AJ007829) Nptl [Cloning vector pGreen]	24	40	2.20E-08	132	82	g3402816	1-194	Bt1G4650	Bt1Gc7716	2171
	TRNA LIGASE) (ASPRS) [Bacillus subtilis]										
	ASPARTYL-TRNA SYNTHETASE (ASPARTATE	∞	52	1.50E-06	121	99	g3122885	147-1	Bt1G4649	Bt1Gc7713	2170
	[Bacillus subtilis]										
	TRANSCRIPTIONAL REGULATORY PROTEIN PHOP										
	ALKALINE PHOSPHATASE SYNTHESIS	52	33	2.50E-12	165	122	g400783	1-383	Bt1G4648	2169 Bt1Gc7708 Bt1G4648	216
	POLYMERASE BETA' SUBUNIT) []									,	
	(TRANSCRIPTASE BETA' CHAIN) (RNA							•			
	DNA-DIRECTED RNA POLYMERASE BETA' CHAIN	6	4	2.90E-08	140	116	g2500611	184-1	Bt1G4647	8 Bt1Gc7707	2168
• -42		2	8 1	1.00E-24	282	439	g460277	101-479	Bt1G4646	7 Bt1Gc7699	2167
	(Y10927) glutamate racemase [Bacillus cereus]	47	36	2.80E-13	174	95	g2462097	413-52	Bt1G4645	5 Bt1Gc7693	2166
	(Z99112) similar to hypothetical proteins [Bacillus subtilis]	99	40	4.90E-23	266	221	g2633912	446-2	Bt1G4644	5 Bt1Gc7677	2165
	(Z99112) similar to hypothetical proteins [Bacillus subtilis]	37	37	6.00E-11	154	131	g2634069	1-296	Bt1G4643	Bt1Gc7674	2164
	PROTEIN) [Bacillus stearothermophilus]				٠						
?.		Cvrg	Ident	Prob	Score	Score			-		NO E
	NCBI of description	%	%	BlastP-	BlastP	aat_	NCBI gi	Position	Gene Id	Contig Id	SEQ

Table 1

MECB [Bacillus subtilis] 16 HYPOTHETICAL HELICASE IN SINI-GCVT INTERGENIC REGION [Bacillus subtilis]	39 1	8.90E-08	132	110	g1731040	268-1	Bt1Gc7956
		3.50E-14 9.00E-08	134	115	g1/2300/	123-1	2200 BUGC/934 BUG4683
Bacill		201	3	3	21722607	1 361	D+10-7034
stearothermophilus] 29 (Z99111) similar to ABC transporter (ATP-binding protein)	73 2	2.70E-59	582	554	g2633814.	474-1	2199 Bt1Gc7916 Bt1G4681
58 50S RIBOSOMAL PROTEIN L20 [Bacillus	70 5	1.50E-21	252	245	g132760	1-210	2198 Bt1Gc7879 Bt1G4680
PROTEIN) (FLAVOHEMOGLOBIN) [Bacillus subtilis] 37 (Z99113) similar to UTP-glucose-1-phosphate	27 3	1.60E-06	116	88	g2634201	333-1	2197 Bt1Gc7863 Bt1G4679
AMMONIA CHAIN) [Bacillus subtilis] 15 FLAVOHEMOPROTEIN (HAEMOGLOBIN-LIKE	64 1	6.50E-14	186	170	g1708267	372-547	2196 Bt1Gc7868 Bt1G4678
PYRIMIDINE-SPECIFIC, LARGE CHAIN (CARBAMOYL-PHOSPHATE SYNTHETASE							
SPOIR-GLYC INTERGENIC REGION [Bacillus subtilis] 5 CARBAMOYL-PHOSPHATE SYNTHASE,	49	5.60E-07	128	81	g115626	120-258	2195 Bt1Gc7854 Bt1G4677
	78 56	5.40E-33	360	353	g732387	1-254	2194 Bt1Gc7858 Bt1G4676
thuringiensis] 4 (AJ004803) putative undecaprenyl-phosphate N	45 34	2.20E-20	241	188	g2582651	1-387	2193 Bt1Gc7856 Bt1G4675
subtilis] 8 IMMUNE INHIBITOR A PRECURSOR [Bacillus	62	1.50E-10	159	137	g124464	1-164	2192 Bt1Gc7853 Bt1G4674
		4.90E-11	163	151	g1709188	1-269	Bt1Gc7852
16 ORF 13231C [Dacillus uluringlensis]	44 17	2.50E-07	9 6 124	100 ¥	g2632519	163-1	2190 Bt1Gc7836 Bt1G4670
subtilis		0 00022	00	2	2226660	304	B+1C-7828
	42 30	8.20E-43	457	390	g585648	834-1	2188 Bt1Gc7823 Bt1G4668
		7.40E-13	170	228	g2633216	428-79	
	35 28	5.00E-11	159	131	g132246	1-314	2186 Bt1Gc7814 Bt1G4666
15 STAGE V SPORULATION PROTEIN D (SPORULATION SPECIFIC PENICILLIN-BINDING PROTEIN) [Bacillus	46 1	7.50E-16	208	185	g586022	285-1	2185 Bt1Gc7767 Bt1G4665
NCBI gi description	% % Ident Cvrg	BlastP- o	BlastP Score	aat_ nap Score	NCBI gi	Position	SEQ ID Contig Id Gene Id NO

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2223 Bt1Gc8106 Bt1G4707 1-383 g	2222 Bt1Gc8102 Bt1G4705 1-179 g	Bt1Gc8099 Bt1G4704 310-1	2219 Bt1Gc8083 Bt1G4703 253-1 g	2218 BITG680/6 BITG4/02 200-21 B	B+1C-8076 B+1C/703 200 21	2217 Bt1Gc8074 Bt1G4701 164-1 g		Bt1Gc8071 Bt1G4700 289-1	Bt1Gc8059 Bt1G4699 19-267	2214 Bt1Gc8067 Bt1G4698 1-168 g		2213 Bt1Gc8064 Bt1G4697 335-1 g	2212 BITOCOOTO BITOTOSO 1-200 B	D+1C-90/19 D+1C/406 1 296	2211 Bt1Gc8055 Bt1G4695 228-1 g	Bt1Gc8044 Bt1G4694 443-1		2209 Bt1Gc8024 Bt1G4692 1-388 g		2208 Bt1Gc8013 Bt1G4691 67-468 g			2206 BITUC8000 BITU4689 62-332 g	Bt1Gc/990 Bt1G4688 1-212	Bt1Gc7965 Bt1G4687 78-239		2203 Bt1Gc7954 Bt1G4686 61-312 g/	NO COURT IN CENETA LOSINON
g732327	g3132723 g730399	g226781	g2851553	8/1/1/9	117700	g1929340		g417115	g2633523	£2632231		g586022	81004/0	122476	g4416482	g2633912		g1770066		g3183562	g755588		g2143613	g3183561	g2632453		g321919	MCDI 61
·355	159	68	78	<u>‡</u>	1	104		187	166	109		176	212	7	101	203		138		148	247		1/8	174	179		103	nap s
384	169	125	112	10/	167	126		200	176	130		182	270		115	230		160		195	284	791,	197	219	188		102	Score
1.50E-35	2./0E-0/ 1.10E-11	1.60E-06	1.30E-05	1.50E-12	1 600 13	4.10E-07		2.90E-15	1.70E-13	1.30E-08		4.70E-13	2.00E-20	2 ME 26	2.40E-06	3.20E-19		9.00E-11		5.90E-15	1.00E-23		1.00E-15	4.70E-18	9.10E-15		1.20E-05	Prob
60	47	35	36	S	h	45		42	5 1	45		37	8	2	<u>သ</u>	39		33		43	47		4/	: £	65	:	35	Ident C
41			15	42		9				70		17	ď		24			22		30	13	<u>.</u> _	99			_	56 1	Cyrg
PROBABLE 1,4-DIHYDROXY-2-NAPHTHOATE	(AF00394) enolase [Stapnylococcus aureus] PRKA PROTEIN [Bacillus subtilis]	RNA polymerase beta' [Pseudomonas putida]	PROBABLE PHOSPHOMANNOMUTASE (PMM)	nypometical protein B - Clostridium acetooutylicum [Clostridium acetobutylicum]	[Bacillus subtilis]	(Z93767) alpha-acetolactate synthase protein, AlsS	subtilis]	PROTOPORPHYRINOGEN OXIDASE (PPO) [Bacillus	(Z99110) similar to thiamin biosynthesis [Bacillus subtilis]	(AJ222587) YkuJ protein [Bacillus subtilis]	SPECIFIC PENICILLIN-BINDING PROTEIN) [Bacillus cubrilici	STAGE V SPORULATION PROTEIN D (SPORULATION	KNA FOLTMEKASE SIOMA-O FACTOR (STAGE III SPORTI ATTON PROTEIN G) [Bacillus subtilis]	DATA DOI VALCE ASE SIGMA OF ASECTOR (STACE III	(AF125999) daunorubicin resistance protein A	(Z99112) similar to hypothetical proteins [Bacillus subtilis]	subtilis]	(Z75208) acetolactate synthase large subunit [Bacillus	ALPHA) (VEGETATIVE PROTEIN 63) (VEG63)	SUCCINYL-COA SYNTHETASE BETA CHAIN (SCS-	13 (U23149) DNA polymerase [Bacillus stearothermophilus]	(magment) - Listeria monocytogenes [Listeria	5'-phosphoribosyl-glycinamide synthetase (EC 6.3.4.13)	VEGETATIVE PROTEIN 296 (VEG296) [Bacillus subtilis]	(Z99104) ybcf [Bacillus subtilis]	plasmid NTP16 []	hypothetical 16.9K protein - Salmonella typhimurium	MCDI gi description

Table 1

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2241	2239 Bt1Gc8243 2240 Bt1Gc8245	2238 Bt1Gc8242	2237 Bt1Gc8231	2236	2235	2234	2233		2231	2230	2229	2228		2227 Bt1Gc8130 Bt1G4711		2226	2225	2224	C	E SEQ	,
BtlG	Bt1Gc8243 Bt1Gc8245	Bt1G	Bt1G	Bt1Gc8230	Bt1Gc8222	Bt1Gc8199	Bt1Gc8186	Bt1Gc8178	Bt1Gc8177	Bt1Gc8172	Bt1Gc8140	Bt1Gc8133		Bt1G		Bt1Gc8120	Bt1Gc8124	Bt1Gc8109	:	Cont	
c824	c824.	c824:	c823	c823	c822	c819	c818	c817	c817	c817:	c814	c813:		c813		c812	c812	c810		Contig Id	
4 Bt														0 Bt							
Bt1Gc8244 Bt1G4728	Bt1G4726 Bt1G4727	Bt1G4725	Bt1G4724	Bt1G4723	Bt1G4721	Bt1G4720	Bt1G4719	Bt1G4718	Bt1G4717	Bt1G4716	Bt1G4713	Bt1G4712		G47		Bt1G4710	Bt1G4709	Bt1G4708	٠.	Gene Id	
28	26 27	.25	24	23	21	20	19	18	17	16	13	12		11		10	9	80		<u>a</u>	
335-1	280-1 450-1	260-1	206-1	1-550	389-1	1-336	360-1	1-240	293-1	372-432	1-287	223-80		437-1		449-1	377-1	283-1		Position	
5-1	2 2	7	<u> 5</u>	50	7	36	<u>-</u>	40	<u></u>	432	87	-80		7-1		1	7-1	<u>-1</u>		tion	
g2 (<u>82</u>	g4 (g4.5	g1(<u>8</u> 13	g21	823	g72	g19	gíl2	g13	g4.5		g17		g <u>2</u> 8	g35	6 13		Z	
g2635766	g2116761 g2960100	g464794	g4582216	g1075694	g135196	g2127280	g2500058	g729328	g1945649	g 1255196	g1351054	g4512408		g1710618		g2829488	g399891	g1351856		NCBI gi	
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183	103 124	204	227	349	224	123	137	269	145	109	106	241		%		300	152	114		7 F	
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1.70E-13	7.50E-05 9.10E-08	1.20E-15	7.70E-18	1.60E-31	4.80E-18	2.40E-08	3.80E-08	2.40E-23	3.30E-10	2.80E-05	9.20E-05	2.20E-20		0.0041		1.20E-26	5.90E-11	1.40E-05		ĕ Ģ	į
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41	37 22	4 8	67	္ထ	39	27	34	2	32	90	27	92		40		47	31	35		% % %	
26 (23 (S	-	10 (34 F	31]	30 F	16 I	31 (ა () [=	51 (<u> </u>	46 F	⊾ N	35 (60 /	5		rg °	
meerculosis] (Z99120) similar to hypothetical proteins [Bacillus subtilis]	subtilis] (D86418) YfnC [Bacillus subtilis] (AL022121) hypothetical protein Rv3676 [Mycobacterium	aureusj STAGE IV SPORULATION PROTEIN A [Bacillus	(AJ237696) elongation factor G (EF-G) [Staphylococcus	LIGASE) (TYRRS) [Bacillus caldotenax] pheromone cAD1 binding protein precursor - Enterococcus from it = 10 mid = 10 mi	[Bacillus thuringiensis] TYROSYL-TRNA SYNTHETASE (TYROSINETRNA	histidine protein kinase (Tn5401) - Bacillus thuringiensis	FORMATE ACETYLTRANSFERASE (PYRUVATE	GLUC	(Z94043) hypothetical protein [Bacillus subtilis]	[Listeria monocytogenes] (U50744) BSMA [Bacillus stearothermophilus]	PREPROTEIN TRANSLOCASE SECA SUBUNIT	(AB017508) rpsS homologue (identity of 87% to B. subtilis) [Bacillus halodurans]	tuberculosis]	PROBABLE RIBONUCLEASE PH (RNASE PH) (TRNA	2 (GSA) (GLUTAMATE-1-SEMIALDEHYDE AMINOTRANSFERASE) (GSA-AT) [Bacillus subtilis]	lactis]	ATP PHOSPHORIBOSYLTRANSFERASE [Lact	OCTAPRENYLTRANSFERASE (DHNA- OCTAPRENYLTRANSFERASE) [Bacillus subtilis] ACONITATE HYDRATASE, CYTOPLASMIC (CIT			
20) s	118) 2212	SEIV Ses	18 pia	SE) (lus the	ine p	A II	COSE	43) h	rıa m 744) I	ROT	1750) cillus	ulosi	ABI	ŠŠ ŠŠ	[AM	SOHG	PRE			
sj imila	18) YfnC [Bacillus subtilis] 18) YfnC [Bacillus subtilis]	SPO) eloi	CAL	-TR	otein	AC	OSE 1-DEHYDROGENASE [Bacillus megaterium]	ypot	onoc;	EIN	AB017508) rpsS homo [Bacillus halodurans]	EO HD I L'INANSFERASE) [MYCOORCEFIUM Mosis]	ERI	LUI LANS	ATE-	HOSPHORIBOSYLTRANSFERASE [Lactococcus	TE H	, _		
r to h	[Bar pothe	RUI	pAL ngatio	RS)	giensi VA S	kina	ETY	EHY	hetica	ytoge A [B	[RA]	S hoi durar	7	BON	FER.	1-SE	RIB(TRA		Z	
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otein	v367	EIN	6	recu	YT)	Pasi Bacil	SE	[Ba	illus	lermo	ECA	tity o		(Z))E 2,	FER.	OP Bac		ption	
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tilis]	ium		Su	cus	Ā	S.		<u>m</u>				otilis		N S	Ľ	lactis] GLUTAMATE-1-SEMIALDEHYDE 2,1-AMINOMUTASE	cus	PRENYLIKANSFERASE (DHNA- PRENYLTRANSFERASE) [Bacillus subtilis] ITATE HYDRATASE, CYTOPLASMIC (CITRATE			
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Table 1

24 AMINO-ACID FERMEASE ROCC [Bacillus submits] 25 (X99545) uracil phosphoribosyltransferase [Bacillus	6 , 8	1.90E-13	113	- - 	g/30000 g1765902	160-1	2263 Bt1Gc8420 Bt1G4751
		2 00E-15	3) (၀	730600	1_340	
7 ACONITATE HYDRATASE (CITRATE HYDRO-LYASE)	56	5.10E-11	165	115	g2506131	176-1	2261 Bt1Gc8391 Bt1G4749
elegans] 34 COMF OPERON PROTEIN 1 [Bacillus subtilis]	34	4.00E-12	171	132	g729157	1-475	2260 Bt1Gc8397 Bt1G4748
31 (U88315) similar to acetyltransferases [Caenorhabditis	30	3.80E-08	134	53	g1825778	412-1	2259 Bt1Gc8392 Bt1G4747
INTERGENIC REGION [Bacillus subtilis]							
29 HYPOTHETICAL 36.8 KD PROTEIN IN PHOB-GROES	43	1.50E-13	181	163	g3025119	1-306	2258 Bt1Gc8390 Bt1G4746
MECB [Bacillus subtilis]			٠				
20 NEGATIVE REGULATOR OF GENETIC COMPETENCE	39	3.50E-14	194	119	g586900	1-485	2257 Bt1Gc8372 Bt1G4745
		1:00	į		610101		
		2 SOF-08	130	چ ک	91346148	366-1	Bt1Gc8368
17 (AF008220) amino acid transporter [Bacillus subtilis]	53	5.90E-10	151	209	g2293166	436-193	2255 Bt1Gc8358 Bt1G4743
SYNTHETASE) [Bacillus subtilis]							
(GLUTAMINE AMIDOTRANSFERASE) (GMP							
13 GMP SYNTHASE (GLUTAMINE-HYDROLYZING)		2.50E-06	118	180	g3123227	1-205	2254 Bt1Gc8353 Bt1G4742
41 (D86418) YfnB [Bacillus subtilis]	36	9.90E-09	133	112	g2116760	1-293	2253 Bt1Gc8347 Bt1G4740
CYTOCHROME B/C SUBUNIT [Bacillus subtilis]	-						
49 MENAQUINOL-CYTOCHROME C REDUCTASE	38	3.90E-14	182	154	g1168649	382-1	2252 Bt1Gc8335 Bt1G4739
19 (AJ011676) DNA ligase [Bacillus stearothermophilus]		1.00E-15	207	177	g3688229	386-1	2251 Bt1Gc8318 Bt1G4738
22 ATP-DEPENDENT HELICASE PCRA []	37	2.70E-13	185	168	g3024353	1-463	2250 Bt1Gc8319 Bt1G4737
[Burkholderia sp.]							
14 (U68411) 4-methyl-5-nitrocatechol oxygenase	40	7.30E-09	142	111	g1575605	388-155	2249 Bt1Gc8308 Bt1G4736
11 (J01829) unknown protein [Transposon Tn10]	61	1.70E-06	118	108	g1196998	154-22	2248 Bt1Gc8286 Bt1G4735
[Bacillus subtilis]							
(version 2) - Bacillus subtilis [] 16 (Z99120) similar to 3-hydroxyacyl-CoA dehydrogenase	39	1.30E-08	142	165	g2635780	1-404	2247 Bt1Gc8290 Bt1G4734
8 UNA-directed UNA polymerase (EC 2.7.7.7) III alpha cham	40	3.10E-10	100	133	g482286	333-1	2246 BITG68280 BITG4/33
		0.001	, 93	161	g2619056	124-256	
))	3				
15 (Z99120) similar to butyryl-CoA dehydrogenase [Bacillus	53	2.20E-15	203	168	g2635778	1-279	2244 Bt1Gc8263 Bt1G4731
37 (Z92952) ywqM [Bacillus subtilis]		9.80E-18	216	199	g1894751	322-1	2243 Bt1Gc8258 Bt1G4730
18 IMMUNE INHIBITOR A PRECURSOR [Bacillus	78	9.80E-50	518	475	g124464	423-45	2242 Bt1Gc8252 Bt1G4729
	Ident Cvrg	Prob	Score	Score	0	-	0
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43 2 2 23	∞ 31	1.20E-08	136	168	24120072	1-0.4		2281 Bt10	
93 22 28					CL99C1V ²) 1	Bt1Gc8509 Bt1G4772		
22 23	1 43	2.00E-21		139	g586058	1-434	Bt1Gc8484 Bt1G4771	2280 Bt10	
93	6 46	1.10E-06	116	97	g2337805	166-1	Bt1Gc8497 Bt1G4770	2279 Bt10	
	5 98	2.80E-75		821	g96497	1-493	Bt1Gc8491 Bt1G4769	2278 Bt10	
40 (AB000617) YceH [Bacillus subtilis]	3 42	2.40E-23		218	g2415745	1-436	Bt1Gc8480 Bt1G4768		
ω	_	4.70E-24	296	252	g2342601	458-1	Bt1Gc8476 Bt1G4767		
[Corynebacterium glutamicum]									
gene; cDNA EST E [] 16 HYPOTHETICAL 69.1 KD PROTEIN (ORF4)	0 80	8.50E-40	427	419	g1730878	1-317	Bt1Gc8475 Bt1G4766	2275 Bt10	
from this gene; cDNA EST EMBL:D32723 comes from this									
comes from this gene; cDNA EST EMBL:D32335 comes									
ankyrin (PIR Acc. No. S37771); cDNA EST EMBL:T01923									
6 (Z703	2 34	2.20E-12	183	122	g3879121	1-434	3c8468 Bt1G4765	2274 Bt1Gc8468	
								٠.	
COA LIGASE) (ACYL-ACTIVATING ENZYME)									
23	1 40	2.30E-21	258	217	g728788	393-1	Bt1Gc8470 Bt1G4764	2273 Bt10	
67 (6 29	5.70E-06	106	88	g2293156	377-1	Bt1Gc8457 Bt1G4762	2272 Bt10	
[Schizosaccharomyces pombe]									
27	9 37	1.10E-19	241	177	g3192023	571-1	Bt1Gc8439 Bt1G4760	2271 Bt10	
LIGASE) (TYRRS) []				. ~					
31		7.70E-05	103	175	g135197	1-387	Bt1Gc8444 Bt1G4758	2270 Bt10	
5	6 32	3.00E-06	124	63	g710552	361-1	Bt1Gc8438 Bt1G4757	2269 Bt10	
cereus]									
29 (AF067645) spore germination protein GerIC [Bacillus	3 48	7.90E-13	175	204	g3290177	317-1	Bt1Gc8435 Bt1G4756	2268 Bt10	
. 32	8 41	0.88	: 59	121	g2506915	350-109	Bt1Gc8431 Bt1G4755	2267 Bt10	
								; ·	
13	1 97	3.20E-81	822	820	g4587971	514-1	Bt1Gc8433 Bt1G4754	2266 Bt10	
					(
23 (AF000309) putative serine/threonine kinase	2 33	2.20E-12	176	122	g2209087	555-1	Bt1Gc8427 Bt1G4753	2265 Bt10	
31 HYPC	9 65	8.00E-39	415	440	g1177011	1-405	Bt1Gc8417 Bt1G4752	2264 Bt10	
caldolyticus									
% NCBI gi description	% % Ident Cvrg	BlastP- Prob	Score Score	nap Score	NCBI gi	Position	Contig Id Gene Id		
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(version 2) - Bacillus subtilis [] PROBABLE FERRIC REDUCTASE TRANSMEMBRANE	30	28	5.70E-18	228	124	g2498389	1-659	2301 Bt1Gc8654 Bt1G4793	
DNA-directed DNA polymerase (EC 2.7.7.7) III alpha chain	9	40	1.30E-16	220	163	g482286	406-1	2300 Bt1Gc8646 Bt1G4792	
BETA SUBUNIT) [Bacillus subtilis] (U61226) RfbB [Leptospira interrogans]	41	40	1.10E-20	244	221	g1666507	428-1	2299 Bt1Gc8627 Bt1G4791	
(TRANSCRIPTASE BETA CHAIN) (RNA POLYMERASE						1			
DNA-DIRECTED RNA POLYMERASE BETA CHAIN	=	42	2.40E-15	207	138	g585920	397-1	2298 Bt1Gc8619 Bt1G4790	
ATP SYNTHASE B CHAIN [Bacillus subtilis]	41	40	2.60E-08	127	103	g584815	1-208	2297 Bt1Gc8617 Bt1G4789	
SYNTHASE II (FGAM SYNTHASE II) [Bacillus subtilis]									
subtilis] PHOSPHORIBOSYLFORMYLGLYCINAMIDINE	14	31	1.30E-07	132	97	g131646	304-1	2296 Bt1Gc8610 Bt1G4788	
(AF008220) putative morphine dehydrogenase [Bacillus	35	4	4.60E-20	238	218	g2293328	297-1	2295 Bt1Gc8594 Bt1G4787	
(AF008220) YtpQ [Bacillus subtilis]	22	61	9.10E-15	188	187	g2293212	1-181	Bt1Gc8589	
(version 1) - Bacillus subtilis []									
DNA-directed DNA polymerase (EC 2.7.7.7) III alpha chain	7	34	4.60E-09	149	111	g80261	304-1	2293 Bt1Gc8590 Bt1G4785	
SYNTHETASE [Drosonhila melanogaster]						(
MULTIFUNCTIONAL AMINOACYL-TRNA	c c	26	8.30E-08	138	70	g135103	408-1	2292 Bt1Gc8580 Bt1G4784	
FORMAMIDOPYRIMIDINE-DNA GLYCUSYLASE	34	ŭ O	8.10E-06	601	133	g1204/4	1-087	2291 Bt1Gc85/6 Bt1G4/83	
IKAMI PROTEIN [ESCRETIONIA COII]	: 6	2, 4	1.20E-12	100	100	8404933	1-105	D1100077	
TRAM DECTEDIT [Escharistic soli]	<u>ဒ</u>	3	1 205 12	. 160	160	~464023	1 104	D+1C-9575	
(GLUTAMINE AMIDOTRANSFERASE) (GMP						•			
GMP SYNTHASE (GLUTAMINE-HYDROLYZING)	31	60	5.60E-47	492	487	g3123227	1-482	2289 Bt1Gc8574 Bt1G4781	
cerevisiae						<i>;</i>			
INFORMATION REGULATOR 2) [Saccharomyces	. ;	i		į	į	0	,		
REGULATORY PROTEIN SIR2 (SILENT	23	42	4.90E-22	264	149	g134506	1-500	2288 Bt1Gc8568 Bt1G4780	
(AL023589) hypothetical protein [Schizosaccharomyces	34	29	6.60E-14	187	98	g3135999	561-1	2287 Bt1Gc8567 Bt1G4779	
[Schizosaccharomyces pombe] (AJ236899) hypothetical protein [Streptococcus gordonii]	41	40	2.40E-07	118	95	g4584073	1-206	2286 Bt1Gc8552 Bt1G4778	
PUTATIVE TRANSPORTER C11D3.18C	. 35	22	3.80E-13	181	112	g1351714	576-1	2285 Bt1Gc8541 Bt1G4776	
[Bacillus subtilis]	,)		,)	, ,) 		
burgdorteri] (AB001488) PROBABLE DNA TOPOISOMERASE III	19	42	8.50E-15	199	172	g1881236	417-1	2284 Bt1Gc8531 Bt1G4775	
(AE001145) prolyl-tRNA synthetase (proS) [Borrelia	14	39	1.80E-06	119	8	g2688299	1-207	2283 Bt1Gc8540 Bt1G4774	
Franciscon Priori	Cvrg	Ident	Prob	Score	Score		* 0311011		
NCRI ai description	%	%	BlastP-	BlastP	88t_	NCRI ei	Position	SEQ Contig Id Gene Id	
			1						

2310 Bt1Gc8692 Bt1G4802	2309 Bt1Gc8679	2308 Bt1Gc8675	2307 Bt1Gc8670	2306 BITGC8664 BITG4/98		2305 Bt1Gc8662 Bt1G4797		2304 Bt1Gc8656		2303 Bt1Gc8660 Bt1G4795	2302 Bt1Gc8641 Bt1G4794		SEQ ID Contig Id NO
692				004		662		656		660	641		Id
Bt1G4802	Bt1G4801	Bt1G4800	Bt1G4799	Bt1G4/98		Bt1G4797		Bt1G4796		Bt1G4795	Bt1G4794		Gene Id
1-374	353-32	349-1	. 1-206	3-431		393-1		324-1		1-409	1-200		Position
g2633908	g1168595	g2739435	g2226222	g20/2/20		g729934		g115950		g282384	g2634202		NCBI gi
115	71	232	72	111		. 68		.204		132	186		nap Score
150	110	267	108	158		117		269		125	189		BlastP Score
6.40E-10	6.90E-06	4.30E-22	1.10E-05	1.20E-10	 } }.	3.00E-07		2.40E-23		2.10E-07	7.10E-15		BlastP- Prob
3 4	35	49	34	32	}	37		51		29	53	-	% % Ident Cyrg
29	38	16	26	29		57		32		41	35		%
stearothermophilus] 29 (Z99112) similar to acetylomithine deacetylase [Bacillus subtilis]	38 ATP SYNTHASE GAMMA CHAIN [Bacillus	(U86377) (p)ppGpp synthetase [Bacillus subtilis]	26 (Y14082) hypothetical protein [Bacillus subtilis]	29 (295121) hypothetical protein Rv3253c [Mycobacterium hiberculosis]	I) [Bacillus caldolyticus]	57 SIGNAL PEPTIDASE I (SPASE I) (LEADER PEPTIDASE	(CATABOLITE CONTROL PROTEIN) [Bacillus subtilis]	32 GLUCOSE-RESISTANCE AMYLASE REGULATOR	subtilis]	41 quinol oxidase aa3-600 chaze II - Bacillus subtilis [Bacillus	(Z99113) similar to alkaline phosphatase [Bacillus subtilis]	COMPONENT [Candida albicans]	NCBI gi description

SEQID NO:

A sequential SEQ ID NO: is assigned to each contig or singleton and the SEQ ID NO: corresponds to that set forth in the sequence listing.

Contig ID

Contigs or singletons are assigned an arbitrary contig ID. Contigs are assembled according to the procedure set forth in Example 2.

Gene ID

Refers to an arbitrarily assigned Gene ID number:

Position

the complement of the sequence set forth in the sequence listing. In cases where the first numeral is higher than its corresponding second numeral, the B. thuringiensis protein or fragment thereof is encoded by forms part of the codon that encodes the C-most terminal amino acid of the coding sequence of the B. thuringiensis protein or fragment thereof. numeral under the position heading is higher than that found in the corresponding second position it designates the nucleotide position which the codon that encodes the N-most terminal amino acid of the coding sequence of the B. thuringiensis protein or fragment thereof. If the first If the first numeral under the position heading is lower than the second numeral, it designates the nucleotide position which forms part of

Each sequence in the GenBank public database is arbitrarily assigned a unique NCBI gi (National Center for Biotechnology Information GenBank Identifier) number. In this table, the NCBI gi number which is associated (in the same row) with a given contig or singleton refers to the particular GenBank sequence which is the best match for that sequence.

aat nap score

based on the BLOSUM62 scoring matrix. The aat nap score is reported by the nap program in the aat package. It is an alignment score in which each match and mismatch is scored

Blastp-Prob

The entries in the "Blastp-Prob" column refer to the probability that such matches occur by chance

BlastP Score

designated clone with the designated GenBank sequence. Each entry in the "BlastP Score" column of the table refers to the BLASTP score that is generated by sequence comparison of the

% Iden

length of that portion of the sequences which is aligned by the BLAST comparison to generate the statistical scores presented The entries in the "%Iden" column of the table refer to the percentage of identically matched nucleotides (or residues) that exist along the

% cvrg <u>100).</u> The %coverage is the percent of hit sequence length that matches to the query sequence (%coverage = (match length / hit total length) x

NCBI gi description

The "NCBIgidesc" column provides a description of the NCBIgi referenced in the "NCBIgi" column.

The following tables, Table 2 through Table 5, are offered by way of illustration and not by way of limitation. It is to be understood that the present invention is not limited to the particular proteins or polypeptides or particuler coding nucleotide sequences listed in Table 2 through 5.

Table 2. Sigma Factor Homologs

MCBI & describnon	м %	% tnahl	-41281H Gorq	YISBIN Score	188 ARA	NCBI &	Gene 1d	ID NO
•	8140	Ident	0011	Score	qsa Score			ON AT
RNA POLYMERASE SIGMA-H		67	1.80E-13	113	Score	g548832	BÜGZZZ	≯81 _{.1€}
FACTOR (SIGMA-30) [Pseudomonas			•					
SIGWY-K EVCLOK bKOCE22ING sem8inoss]	66	*	1.30E-13	LLI	ELI	ETEIPIB	Buchii	331
REGULATORY PROTEIN BOFA								
(BYPASS-OF-FORESPORE PROTEIN)								
KNA POLYMERASE SIGMA-G [Bacillus subtilis]		68	7.30E-119	SLII	1164	2748813	BilG627	66Þ
PACTOR (STAGE III SPORULATION								
PROTEIN G) [Bacillus subtilis]		70	1L 200 V	012	33L	0000010	のしろいけは	OUF
FACTOR PRECURSOR [Bacillus		96	4.90E-71	61 <i>L</i>	SSL	8133289	Bt1G628	66Þ
thuringiensis] RNA POLYMERASE SIGMA-28 EACTOR REFILESOR (Recilling	100	100	1.00E-118	6911	6911	£133282	Bt1G1336	9/01
TACTOR PRECURSOR [Bacillus thuringiensis]								
transcription initiation factor sigma H - Bacillus megaterium [Bacillus		9 <i>L</i>	4 70E-66	-ZL9 .	6Ĕ9	L9E7873	BilG1827	1108
megaterium] RNA POLYMERASE SIGMA FACTOR		1/8	9.60E-130	£L71	977 l	9975513	Beigzisz	6 <i>L</i> 11
RPOD (SIGMA-A3) [Bacillus			061 500:5	C / C T		001.55-9	46150115	
SPORULATION SIGMA-E FACTOR subtilis]		85	8.20E-07	611	<i>L</i> 91	£9/7E[3	Bf1G2201	9811
PROCESSING PEPTIDASE (STAGE II		0.0	10-50510	(11	407	6011.679	10770110	(017
SPORULATION PROTEIN GA) [Bacillus		,		,				
[silitdus]		CL	ו פטבייוט	121	121	CIOLECES	Periodon	3061
(STAGE II SPORULATION PROTEIN		7L	J.60E-40	124	431	716/07CA	Bt1G5594	CU21
[Bacillus coagulans]		6 <i>L</i>	1 80F-57	102	105	LSLVELO	Perions	SUCI
SPORULATION PROTEIN AB) [Bacillus		<i>(</i> 1	I.80E-57	169	165		B11G5595	C071
RNA POLYMERASE SIGMA-F FACTOR		83	3.00E-103	1023	1401	069t9t ^g	B41G2296	1205
(STAGE II SPORULATION PROTEIN								
AC) (SPORULATION SIGMA FACTOR)								
[Bacillus megaterium]		ΟV	L> DUB V	LOS	37 _/ 3	, 11,000,00	PHEDIN	LCC I
(Z99111) similar to RNA polymerase sigma factor [Bacillus subtilis]		6 †	4.80E-57	L85	595	01/0023	BilC7414	177 ī
fermanc enmancil towns surface				•			,	

PROTEIM G) [Bacillus subtilis]								•
FACTOR (STAGE III SPORULATION								
thuringiensis]	98	69	7.00E-26	867	7.LT	g133475	B#1C4696	2212
G FACTOR (ORF3) [Bacillus				-		_		
subtilis] POSSIBLE RNA POLYMERASE SIGMA-	IL	0\$	2.30E-16	203	ZLI	1845E1g	Bt1C4171	1 941
RPOD (SIGMA-A) (SIGMA-43) [Bacillus			•		÷			
KNA POLYMERASE SIGMA FACTOR [Streptomyces coelicolor]		33	4.10E-06	114	66	9945E1g	BilG4172	1771
aeruginosa] (ANA polymerase sigma factor (ANI 0320)	18	82	1.90E-14	182	.98	84288958	BI1C3404	1364
FACTOR (SIGMA-30) [Pseudomonas								
SIGV [Bacillus subtilis] RNA POLYMERASE SIGMA-H	66	97	4.50E-13	7 <i>L</i> I	LSI	g548832	Bt1G3335	LSEI
KNA POLYMERASE SIGMA FACTOR [Listeria monocytogenes]	66	52	6.90E-17	208	126	g3024615	BIIC3331	LSEI
gene [Bacillus subtilis] (AF074855) RMA polymerase sigma B	100	53	3.00E-71	127	IIL	e23865g	B#I C3067	1328
FACTOR [Bacillus subtilis] (X93081) sigma F/sigma G transcribed	66	32	7.80E-25	283	240	81941918	B#1@7697	1274
KNA POLYMERASE SIGMA-54 Region [Bacillus subtilis]	100	74	1.90E-85	\$58	. 832	g133292	B#1@7627	1595
WWCE-BEWBAA INTERGENIC			•					
TRANSCRIPTIONAL REGULATOR IN								
PUTATIVE SIGMA L-DEPENDENT	L6	19	8'30E-211	2038	2core 5059	0901£71g	B#1@5233	 1548
255.	Cvrg	Ident	Prob	Score	qsa			ID NO
MCBI gi description	%	%	-Ateala	Alzala	İBB	NCBI gi	Gene Id	ZЕб

SI

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<u>SEQID NO:</u> A sequential SEQ ID NO: is assigned to each contig or singleton and the SEQ ID NO:

corresponds to that set forth in the sequence listing.

Gene D:

Refers to an arbitrarily assigned Gene ID number.

WCBI gi: Each sequence in the GenBank public database is arbitrarily assigned a unique NCBI gi

Wational Center for Biotechnology Information GenBank Identifier) number. In this table, the NCBI gi

mumber which is associated (in the same row) with a given contig or singleton refers to the particular

number which is associated (in the same row) with a given contig or singleton refers to the particular GenBank sequence which is the best match for that sequence.

334 nap score: The sat nap score is reported by the nap program in the sat package. It is an alignment

sat nap score: The sat nap score is reported by the nap program in the sat package. It is an alignmen score in which each match and mismatch is scored based on the BLOSUM62 scoring matrix.

Blastn-Prob: The entries in the "Blastn-Prob" column refer to the probability that such matches occur by

Blastp-Prob: The entries in the "Blastp-Prob" column refer to the probability that such matches occur by chance.

BlastP Score: Fach entry in the "BlastP Score" column of the table refers to the BI ASTP score that is

BlastP Score: Each entry in the "BlastP Score" column of the table refers to the BLASTP score that is generated by sequence comparison of the designated clone with the designated GenBank sequence.

Modern The entries in the "Miden" column of the table refer to the percentage of identically matched mucleotides (or residues) that exist along the length of that portion of the sequences which is aligned by nucleotides (or residues) that exist along the length of that portion of the sequences which is aligned by

the BLAST comparison to generate the statistical scores presented $\frac{\& \text{ cvrg:}}{\& \text{ cvrg:}}$ The %coverage is the percent of hit sequence length that matches to the query sequence (%coverage = (match length / hit total length) x 100).

(%coverage = (match length / hit total length) x 100).

WCBIgi" column.

"NCBIgi" column.

Table 3. Transposases, Integrases, and Transposons

TRANSPOSASE FOR INSERTION	57	c 8	4'40E-21	072	LES	28E7642g	416 Brigs12
Bacillus thuringiensis [Bacillus							
transposase (insersion sequence IS231) -	35	34	I.80E-07	123	LS	0627212g.	410 BHG207
[siznəignirudi							
SEQUENCE ELEMENT IS231E [Bacillus						_	
TRANSPOSASE FOR INSERTION	12	ZL	7.50E-26	967	TEE	g249113	387 Bt1G475
influenzae Rd]						·	
PROTEIN HII721 [Haemophilus		0.0	u innere			00-16-29	0010119 505
HADOLHELICYT LKYNSDOSYSE-FIKE intineuzse kg]	LS	36	3.30E-17	112	171	004764 <u>Cg</u>	383 Bi1G468
PROTEIN HII721 [Haemophilus							
PROTEIN HILZL TRANSPOSASE-LIKE	nο	38	4.10E-19	677	18t	62497400	319 Bt1G464
thuringiensis]	09	86	01 401 1	000	701	0072070-	· ·
SEQUENCE ELEMENT IS231F [Bacillus					•		
TRANSPOSASE FOR INSERTION	61	08	2.70E-38	410	⊅ L€	\$24611d	226 Bt1G276
[Anabaena PCC7120]		•		•			
(AF047044) putative transposase	98	77	4 ′30E-0∂	140	£6	£3005554	94 BilG88
sequence IS232]							
SEQUENCE ELEMENT IS232 [Insertion			٠,			•	
TRANSPOSASE FOR INSERTION	33	34	2.30E-09	SÞI	120	28E794282	2 Biigz
•	9	111000		21022	nap Score	•	
แจกจุ่นเรอา 13 เสวาา		Ident		Score		ig tacki	ID NO SEÓ CEUE IQ
MCBI gi description	%	%	-Aisal	PlastP	100	NCBI gi	SEQ Gene Id

					-			
diringiensis]				-				
SEQUENCE ELEMENT IS231C [Bacillus					c			
TRANSPOSASE FOR INSERTION		66	6.60E-250	7042	2449	· \$136144	BeiG2554	1252
sequence [S232]								
SEQUENCE ELEMENT IS232 [Insertion		"	70-7011	LLC	228	785794582	C+C70.110	L171
LKYN2bO2Y2E ŁOK INZEKŁION sedneuce [2737]	VC	66	1.70E-52	7 75	863	CSELOVCO	Delicosys	VICI
SEQUENCE ELEMENT IS232 [Insertion								
TRANSPOSASE FOR INSERTION		66	6.50E-53	848	232	285794282	מנזמזום	/071
sequence IS232]	30	00	C 201 7	0/3	663	COLLUVE	D+1(5)303	LUCI
SEQUENCE ELEMENT IS232 [Insertion								
TRANSPOSASE FOR INSERTION		9 <i>L</i>	9.10E-26	767	987	2857497382	C1779119	0611
sequence IS232]		,,,)C 401 0	000	,	COCLOVC .	710001,4	0011
SEQUENCE ELEMENT IS232 [Insertion								
TRANSPOSASE FOR INSERTION		7 6	1.20E-44	0 <i>L</i> t	424	2857945282	BilG1818	860 I
[Anabaena PCC7120]								
(AF047044) putative transposase	l þ	74	1.60E-11	195	102	t g3005554	Bugilt	7/01
sedneuce IS232]								
SEQUENCE ELEMENT IS232 [Insertion			•		/			
TRANSPOSASE FOR INSERTION	S †	16	2.60E-88	288	716	788794282	BilClezd	S † 01
influenzae Rd]			S					
PROTEIN HII721 [Haemophilus						_	**	
HYPOTHETICAL TRANSPOSASE-LIKE	LS	0 7	2.50E-19	231	183	004764 <u>6</u> 2	Bt1G1238	1002
thuringiensis]						•		
(Y09946) transposase [Bacillus		91	4'60E-35	321	308	07794718	BriG1481	086
(control de la control de la c						0.00.0019.		
(Y09450) transposase [Pseudomonas	66	30	2.80E-37	111	358	8684691g	Bitelin	00 <i>L</i>
(AE000433) IS150 putative transposase [Escherichia coli]	cc	₽ \$	1.60E-26	567	922	18668718	חחוחות	10/
[71enr nosogener]]	36	VS	3C-309 1	900	9LC	18008712 6	שיוניוטפּ	107
71eaT nosogsnert SCIAq bimasılq								:
transposase mpA - Enterococcus faecalis	₽¢ ·	56	2.70E-25	566	761	716618	BilG913	169
[sbiida]	,,	,,	30 402 0	000	701	CDOOD	CIODIA	.05
(Y09450) transposase [Pseudomonas	LE.	98	I'70E-61	635	L6 S	g1694898	BHIG913	169
sequence IS232]				:				
SEQUENCE ELEMENT IS232 [Insertion								
TRANSPOSASE FOR INSERTION	9	96	3.40E-07	172	601	28£74442g	BII C864	099
[iloo sidoricheal]			•		•			
(AE000433) IS150 putative transposase	70	77	1.10E-08	132	110	1866871 <u>8</u>	Bt1G843	bb 9 .
[sastuboibst								
(AB016803) transposase [Deinococcus	58	87	3.30E-33	362	288	g3426013	Bt1C840	7 1 9
[Anabaena PCC7120]								
(AF047044) putative transposase	82	67	2.60E-09	741	6 <i>L</i>	£300554	Bt1G782	£09
[Anabaena PCC7120]					••			
(AF047044) putative transposase	77	57	4.00E-06	113	98	g3005554	TOTO 118	555
sequence [S232]								
SEQUENCE ELEMENT IS232 [Insertion	ıc	46	CO-3100.4	140	000	2857942g	1400110	OFC
LKYN2bO2Y2E ŁOK INZEKLION sedneuce [2535]	31	76	9.00E-63	149	909	COLLOPCO	ומאטוים	7V3
SEQUENCE ELEMENT 18232 [Insertion								
CEOUTH COURT IS SOMETIONS					Score			
	Cvrg	Ident	Prop	Score			•	ID NO
NCBI gi description	%	%	-Arsala	PlastP		NCBI gi	Gene Id	SЕÓ

MCBI gi description	້ ,‱	% Ident	-Assala dorA	BlastP	taa qan	NCBI gi	Gene Id	ID NO
esesonement evitation (AMOTANTA)					Score	, , , , , , , , , , , , , , , , , , ,	איונטעטוא	
AF047044) putative transposase [Anabaena PCC7120]		74	7.20E-09	138	100		Bi1C7694	
LM224) [Staphylococcus amens]		67	6.30E-71	812	654	9566518	Bt1G2705	<i>\$L7.</i> 1
U66614) putative transposase	100	38	9.90E-41	433	342	Z19860Zg	Bt1C5002	1302
Marinococcus halophilus] [AE000433] IS150 putative transposase	90 (25	7.20E-38	901	785	18668718	Bt1G2979	LIEI
Escherichia coli] [AF047044) putative transposase	35 (52	9.60E-09	136	66	\$2005g	Bt1G3087	1330
TRANSPOSASE FOR INSERTION FOR THE PROPERTY OF	. 001	100	2.90E-231	2231	2231	28£7949	B41G3161	1340
edneuce ISS3S] ZEÓNENCE ETEMENT ISS32 [Insertion						٠.		
SEQUENCE ELEMENT IS231C [Bacillus	S L S I	† 6	7.10E-31	340	916		Bt1G3278	esei
huringiensis] (AB016803) transposase [Deinococcus) † 6	30	6.90E-49	210	330	£3426013	Bt1G3789	1324
adiodurans] TRANSPOSASE FOR INSERTION TRANSPOSASE FOR INSERTION	. 67	\$6	9.30E-61	229	919	285794282	Bt1G3378	1362
ACOUNTY ELECTION SEQUENCE ELEMENT IS232 [Insertion	S .	CJ		800	OSC	OLLOVLI	DelCaeon	
Y09946) transposase [Bacillus :	1	75	I.70E-27	308	087		Bt1G3280	
PROTEIN HILTS! (Hacmophilus HYPOTHETICAL TRANSPOSASE-LIKE	Ī	8 £	4'10E-16	575	<i>L</i> 81	00 5 76573	B41C3232	8 /£I
nfluenzae Rd] Americae Rd]	100 (25	3.00E-80	908	96 <i>L</i>	18668718	Bt1G3625	1380
Escherichia coli] AE000433) ISI50 putative transposase	100 (15	I 40E-80	608	822	18668718	B11C3995	£8£1
Escherichia coli] AF047044) putative transposase) ZL	50	7.90E-17	208	LII	\$3005554	Bt1C3984	1386
Anabaena PCC7120] AL023861) putative IS element	100 (56	2.80E-20	240	69 I	02E812Eg	Bt1G3687	1386
ransposase [Streptomyces coelicolor] AF047044) putative transposase	36 (74	1.50E-08	132	<i>L</i> 6	422005g	Bt1G3704	1388
Anabaena PCC7120] Y09946) transposase [Bacillus	100 (IS	9.90E-73	132 ·	SEL	0779471g	Brig3745	1391
hwringiensis] TRANSPOŠASE FOR INSERTION ""	1001	100	1.00E-251	7474	2443	8136144	Bt1G3762	1392
huringiensis] Hypothetical Transposase Ike	1	86	2308 07	061		0012012	D+1(53043	0/51
PROTEIN HII721 [Hacmophilus PROTEIN HII721 [Hacmophilus	I	97 ⁷⁷²⁵	5'30E-07	150	76	00+/6+78	Bt1G3345	6 6 01
SEQUENCE ELEMENT IS232 [Insertion	717	43	4.60E-13	6 <i>L</i> I	123	285794282	Bt1G3970	ELSI
equence IS232]	S	67	1.00E-08	136	159	285794S	B#1C4000	1191
SEQUENCE ELEMENT IS232 [Insertion	S							

MuA2 [Zea mays]	ī				<u>.</u> .		
hypothetical protein 612 - maize transposon	l þ l	59	1.60E-20	721	235	725974 ₈	2055 Bt1G4494
[AC006217] putative retrotransposon	10	SL	6.20E-46	767	SLV	£9790£43	1888 B41C4413
Tn10] (J01829) unknown protein [Transposon	18	IS	3.00E-13	180	160	8669611 ₃	1197 Bt1G4204
Tn4536] (101829) unknown protein [Transposon		çç	2.50E-16	208	220	8669611g	1795 Bt1G4202
HYPOTHETICAL 37.1 KD PROTEIN IN		32	1.40E-08	9EI	₽ L	8141420	1340 Bt1C3160
Tn1546] CARBOXYPEPTIDASE) [Transposon)						
beblidyse) (DD- Cybroxabeblidyse (DD-		•					•
D-VTVAXF-D-VTVAINE redneuce [2737]	;	<i>L</i> 9 '	1.90E-13	8/1	8 <i>L</i> I	902985g	105 Bile134
SEQUENCE ELEMENT IS232 [Insertion	13.	19	6.20E-14	78 1	130	285794282	7123 B41C4631
sedneuce [2532] SEONENCE ELEMENT [5232 [Insertion	;					-	
TRANSPOSASE FOR INSERTION sequence [S232]	:	64	1.50E-18	575	₽L!	28£7 0 497382	5132 B41C4615
SEQUENCE ELEMENT IS232 [Insertion sequence IS232]	13	95	1.90E-10	122	130	Z8EL67ZB	5130 B41C4607
TRANSPOSASE FOR INSERTION SEQUENCE ELEMENT IS232 [Insertion sequence IS332]		32	5'40E-10	1 \$1	611	79C16b78	2049 BilG4484
SEQUENCE ELEMENT IS231C [Bacillus thuringiensis]	l	3 C	Of BOV C	121	011	COLLOVC	2000 D+1C0080
SEGUENCE FOR INSERTION SEQUENCE IS232]	. 07	99	4.10E-27	LOE .	997	8136144	2041 Bt1G4473
SEQUENCE ELEMENT IS232 [Insertion		98	1.10E-15	503	812	78£ <i>L</i> 6†73	1965 Bri C4386
(AE000433) ISI 50 putative transposase [Escherichia coli]		32	3.80E-25	987	258		1902 Bt1C4312
CEQUENCE ELEMENT IS231F [Bacillus	i						
LKYNZŁOZYZE ŁOK INZEKLION zedneuce [2737]	. 97	87	2.10E-11	191	ε6	\$116\$5g	1869 Bt1G4277
SEQUENCE ELEMENT IS232 [Insertion	. 22	95	1.10E-20	248	213	Z85794 <u>5</u> 24	1804 BIIC4511
sedneuce [2232] SEGNENCE ELEMENT [S232 [Insertion							.
TRANSPOSASE FOR INSERTION TRANSPOSASE FOR INSERTION	1	61	7.20E-24	927	٠, ٢, ٢	28679422	1139 Bile4145
SEQUENCE ELEMENT IS232 [Insertion	ti .	25	6.90E-11	126	SII	Z8£764 <u>Z</u> g	1725 Bt1G4130
sedneuce ISS3S]			/ I - 710C: I	077		70C1CF7 3	7014D11G 0601
TRANSPOSASE FOR INSERTION			1.50E-17	220	Score 177	CSELOPCO	1698 Bt1C4102
MCBI gi description	W Cvrg	% Ident	-Assala Prob	Score 52002		NCBI ^g i	ID NO SEÓ Ceue Iq
					•		

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NCBI gi description	رىرى رىرى	Ident %	-AjsafA dorA´	BlastP Score	tee qea	NCBI ^g i	Gene Id	ID NO
					Score		•	
MUM nosoqarara - maize transposon MuDM	ÞΙ	IS	4'50E-13	184	524	14105128	B11C4460	Z021
(J01829) unknown protein [Transposon	II	19	1.70E-06	811	108	866961 IB	Bt1C4735	2248
[01 _n T								
IALEGEVESE STATE STATES IN A SECONDE STATES OF THE SECONDE STATES	23	7 <i>L</i>	4'90E-29	909	18 5	£8£01718	BilG21	τī
[Bacillus subtilis]								
U77495) putative integrase [Leuconostoc	35	32	1.20E-12	ELI	\$6 .	g4098413	BilG327	967
oenos bacteriophage 10MC]		00	.,		,,,	2000017	1,5001.4	-
(AL035707) putative integrase	100	67	1.10E-41	744	364	L6606+1+3	Bt1G961	77.L
(AB001488) PROBABLE INTEGRASE. [Streptomyces coelicolor]	\$6	67	3.20E-35	185	767	16218812	Bi1G1164	834
[Bacillus subtilis]	30	UC	00 200 0	366	100	031331~	37113174	
(M34832) integrase (int) [Bacteriophage phi-11]	C6	30	S'40E-30	332	351	810013	BilGi165	458
(X98106) integrase [Bacteriophage	100	6ξ	1.40E-71	724	£69	81926326	BilC1256	666
phigle]								

SEQID NO: A sequential SEQ ID NO: is assigned to each contig or singleton and the SEQ ID NO:

corresponds to that set forth in the sequence listing.

number which is associated (in the same row) with a given contig or singleton refers to the particular (National Center for Biotechnology Information GenBank Identifier) number. In this table, the NCBI gi NCBI gi: Each sequence in the GenBank public database is arbitrarily assigned a unique NCBI gi Refers to an arbitrarily assigned Gene ID number.

aat nap score: The aat nap score is reported by the nap program in the aat package. It is an alignment GenBank sequence which is the best match for that sequence.

Blastp-Prob: The entries in the "Blastp-Prob" column refer to the probability that such matches occur by score in which each match and mismatch is scored based on the BLOSUM62 scoring matrix.

generated by sequence comparison of the designated clone with the designated GenBank sequence. BlastP Score: Each entry in the "BlastP Score" column of the table refers to the BLASTP score that is chance.

the BLAST comparison to generate the statistical scores presented nucleotides (or residues) that exist along the length of that portion of the sequences which is aligned by % Iden: The entries in the "Miden" column of the table refer to the percentage of identically matched

% cvrg: The %coverage is the percent of hit sequence length that matches to the query sequence

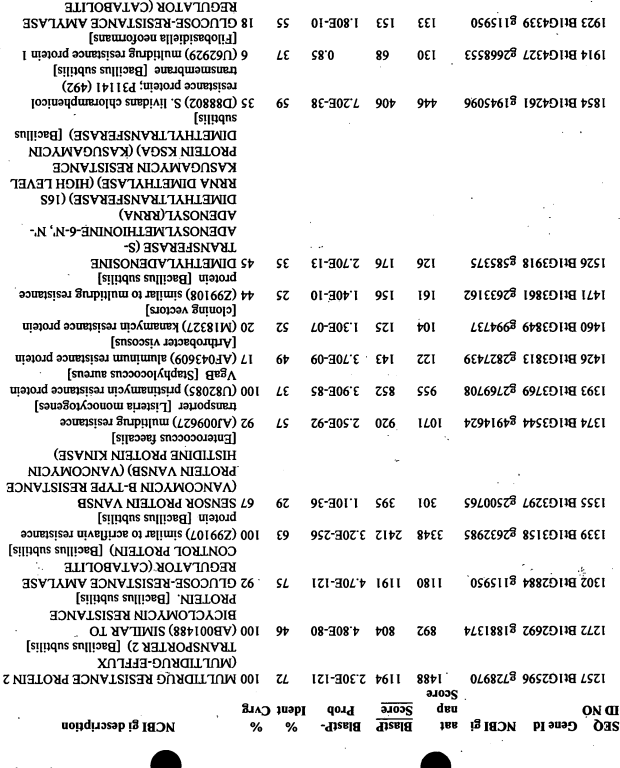
"NCBIgi" column. NCBI gi description: The "NCBIgidese" column provides a description of the NCBIgi referenced in the (%coverage = (match length / hit total length) x = (0).

Table 4. Antibiotic, Chemical, and Heavy Metal Resistance

						Score			
		Cvig	Ident	Prob	Sc re	qвп			ID NO
(CBI gi description	Į	%	%	-ArsalB	BlastP	İBB	NCBI &	Gene Id	ZЕб

MCBI gi description	% %	% Juəbl	BlastP- dor9	BlastP	taa qan	NCBI gi	D NO Gene Iq	
•	•		•		Sc re	2.2//0-		
MULTIDRUG RESISTANCE-LIKE ATP-BINDING PROTEIN MDL [Escherichia	61	56	5'10E-55	£17	520	8159928	33 B41G21	
coli]								
(A1235272) BICYCLOMYCIN	<i>L</i> I	££	4'80E-09	114	†01 :	741198Eg	88 BHC130	
RESISTANCE PROTEIN (bcr1)		•						
IZOFENCAF-LENV ZANLHELVZE' [Kickettzis blowszekii]	Ll	SS	1.30E-49	226	664	9124711g	142 B41G181	
MUPIROCIN RESISTANT	/ T	cc	Characti	070	<i>((</i> L	01CH/119	י -	
(ISOLEUCINE-TRNA LIGASE) (ILERS)								
(MUPIROCIN RESISTANCE PROTEIN)								
[Staphylococcus aureus]					J.,			
noloniup of miliar in mondan (78928X)	95	Lε	4.00E-28	314	435	[8465]	162 Bt1G201	-
resistance protein NorA [Bacillus subtilis] (D88802) S. lividans chloramphenicol	01	7 5	1.70E-14	192	183	81945096	180 Bit Coop	
resistance protein; P31141 (492)	4 1	+c	F1-30/11	761	COI	06004618	180 Bt1G555	
transmembrane [Bacillus subtilis]								
ACETYLTRANSFERASE (TABTOXIN	66	EE _	6.10E-25	784	Z3 2	g136472	704 BII C 746	
RESISTANCE PROTEIN) [Pseudomonas								•
Syringae] (X92868) mercuric resistance operon	LV	IV.	00 301 A		J11 **;	0700010	JJE DHICOO	
(X92868) mercuric resistance operon regulatory protein [Bacillus subtilis]	14	It :	6.10E-09	133	911 🤲	69Z801Zg	775 Bt1G332	
regularity of neimies moral (78928Z)	84	38	2.00E-26	867	383	[204831g	598 BHC339	
resistance protein NorA [Bacillus subtilis]								
(A1235272) BICYCLOMYCIN	09	L Z	4'30E-11	512	66 l	741138Eg	391 Bf1C441	
RESISTANCE PROTEIN (ber1)								
Rickettsia prowazekii] BACITRACIU RESISTANCE PROTEIN	<i>ν</i> δ	05	€ 00E-50	3UY	VUY	9CN2ULID	3917114G 705	
(PUTATIVE UNDECAPRENOL (PUTATIVE UNDECAPRENOL	+0	65	2.90E-59	\$09	1 09	81705428	397 Bt1G486	
(KINASE) [Escherichia coli]	•	• ,	r right					
TELLURITE RESISTANCE PROTEIN	09	97 .	7.10E-12	891	6 6	\$1174634	451 BHG219	
TEHB HOMOLOG [Haemophilus						•		
influenzae Rd]	-0	30) ((U) L	3,0	300	57.0070		
(S9108) similar to multidrug resistance	£8	52	7.60E-26	242	359	2915233	453 B41G255	
protein [Bacillus subtilis] (Z82987) unknown similar to quinolon	100	33	9.00E-55	346	889	[28465]	S79 BUG744	
resistance protein NorA [Bacillus subtilis]	;							
(Z99107) similar to acriflavin resistance	99	Lε	7.30E-96	856	<i>L</i> ZII	82632985	613 Bt1G795	
protein [Bacillus subtilis]	20		U1 40L 1	301		2000070		
(AE001125) acriflavine resistance protein	LZ.	77	1.70E-13	681	183	7208892g	954 BHG800	
(acrB) [Borrelia burgdorferi] bacitracin resistance protein homolog bacA	00	82 .	4.20E-10	ttl	LII	91824128	692 Bt1G915	
- Mycobacterium leprae [Mycobacterium	"	07 ·	01-707*+		/ 1 1	0100-179	CIADIIO 740	
leprae]							•	
SENSOR PROTEIN VANSB	100	28	1.20E-37	† 0†	968	22500765	726 Bt1G970	
(VANCOMYCIN B-TYPE RESISTANCE						.=		
PROTEIN VANSB) (VANCOMYCIN							•	
HISTIDINE PROTEIN KINASE)				•			·	
[Enterococcus faecalis]			20. 200 2					
(AF043609) aluminum resistance protein	100	8 <i>L</i>	7.80E-183	⊅ LLI	⊅ ∠LI	62827439	862 Bt1G1220	
[Arthrobacter viscosus]								

TRANSPORTER 2) [Bacillus subtilis]							•
(MULTIDRUG-EFFLUX							
MULTIDRUG RESISTANCE PROTEIN 2	100	17	6.80E-26	293	180	076827g	1546 Bt1G5250
TRANSPORTER 1) [Bacillus subtilis]			•		•		-
(MULTIDRUG-EFFLUX							
MULTIDRUG RESISTANCE PROTEIN 1	100	73	4'30E-54	<i>9L</i> 7	168	7E3134g	154e B41G5216
RESISTANCE GENE. [Bacillus subtilis]							
CEKEUS ZWITTERMICIN A-					•		
(AB001488) SIMILAR TO BACILLUS	100	34	1.40E-41	Itt ,	392	g1881342	1540 B41 C5467
[silitdus							•
AMP TO THEIR SUBSTRATE. [Bacillus							
DEFENDENT COVALENT BINDING OF							
WHICH ACT VIA AN ATP-							
(AB001488) SIMILAR TO ENZYMES	70	98	2.40E-12	173	6\$ I	g1881228	1227 Bt1G2407
KINASE) [Escherichia coli]							2
(PUTATIVE UNDECAPRENOL				***	67.4	071 60179	001 70 110 0771
BYCILKYCIN KESISTANCE PROTEIN	001	SS	2.70E-54	195	EIL	82420719	1526 Bt1G2406
protein [Bacillus subtilis]				701		0011-0023	
protein [Bacillus subtilis] (Z99113) similar to fosfomycin resistance	00	65	8.40E-44	794	<i>ttt</i>	89175960	1208 Bt1G2308
(S9109) similar to multidrug resistance	001	7 t	3.10E-78	L8L	186	+C+CC078	1190 BilG2211
ubtilis]	UUI	CV .	3 105-78	LBL	120	VEVEENCE	1100 1100 11
DIMETHYLTRANSFERASE) [Bacillus	٠.				• '		•
PROTEIN KSGA) (KASUGAMYCIN							
KASUGAMYCIN RESISTANCE							
KASUS ANGRETHYLASE) (HIGH LEVEL						•	
DIMETHYLTRANSFERASE) (165							
ADENOSYL(RRNA)							
VDENOZATWELHIONINE-6-N', N							
TRANSFERASE (S-							
DIWETHYLADENOSINE	100	0 <i>L</i>	1.60E-106	1024	10 44	8282375	1188 BHC5166
NorA [Synechocystis sp.]							
(D90909) quinolene resistance protein	100	28	1.20E-26	300	415	81652918	1130 B41C1638
transmembrane [Bacillus subtilis]			•				
resistance protein; P31141 (492)						_	-
(D88802) S. lividans chloramphenicol	100	97	7.70E-77	<i>8LL</i>	786	9602491g	1053 BHG1297
monocytogenes]							
operon regulatory protein (MerR) [Listeria		•					
(Y07640) putative mercury resistance	St	TE.	3.30E-10	142	110	2779772	1002 BHG1220
[Staphylococcus aureus]				•			
(MUPIROCIN RESISTANCE PROTEIN)							
(ISOLEUCINE-TRNA LIGASE) (ILERS)							
MUPIROCIN RESISTANT	cc	.0.0		0-01	CLOI	016-1119	662 BHC1250
ISOFENCAF-LKNY SANLHELYSE' [Streptomyces bencetins]	23	.85	1.80E-190	9781	1843	9157/110	005 Brite1520
TRANSMEMBRANE PROTEIN							
TE ANSMEMBE AND PROTEIN	co	52	2.60E-17	717	246	8233400	986 Bri C1495
resistance protein [Cloning vector pFW13]	30	36	LI aus c	CIC	370	301005-	30/15/140 300
(U50978) kanamycin/gentamycin-	7 1 2	ΙÞ	1.80E-34	₽L E	₽ L€	CF1//718	950 BHG1408
PROTEIN VANW [Enterococcus faecalis]	.,	• •	1 000 t	720	, 20	20.2201	007101.4 030
VANCOMYCIV B-TYPE RESISTANCE	100	67	3.70E-34	115	332	91166428	930 BtiG1324
					Sc re		
_	Cvrg	Ident	Prob	Score	qsa		ON (II
MCBI gi descripti n	%	%	-AsealA	Alzal	JBB	NCBI gi	SEQ Gene Id



5'40E-09

9.60E-05

5.00E-13

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30

67

SII

L01

181

101

86

238

2211 Bt1G4695 g4416482

2179 Bt1G4659 g2632985

2151 Bt1G4629 g3925779

24 (AF125999) daunorubicin resistance protein A [Mycobacterium avium]

11 (Z99107) similar to acriflavin resistance

CONTROL PROTEIN) [Bacillus subtilis]

31 (AL034353) putative major facilitator family multi-drug resistance protein [Schizosaccharomyces pombe]

protein [Bacillus subtilis]

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NCI	%	%	-ArsalA	Alzal B	jer	NCBI gi	Gene Id	Č	

SEQ Gene Id MCBI gi aat BlastP BlastP % % MCBI gi description

ID NO nap Score
Score

Score

ARGULATOR (CATABOLITE
CONTROL PROTEIN) [Bacillus subtilis]

SEQID NO: A sequential SEQ ID NO: is assigned to each contig or singleton and the SEQ ID NO: Gene ID:

Gene ID:

Refers to an arbitrarily assigned Gene ID number.

<u>MCBI git</u> Each sequence in the GenBank public database is arbitrarily assigned a unique MCBI git number for Biotechnology Information GenBank Identifier) number. In this table, the MCBI git number which is associated (in the same row) with a given contig or singleton refers to the particular number which is the best match for that sequence.

GenBank sequence which is the best match for that sequence.

<u>ast_nap score</u>: The sat_nap score is reported by the nap program in the sat package. It is an alignment score in which each match and mismatch is scored based on the BLOSUM62 scoring matrix.

<u>Blastp-Prob</u>: The entries in the "Blastp-Prob" column refer to the probability that such matches occur by

chance.

BlastP Score: Each entry in the "BlastP Score" column of the table refers to the BLASTP score that is generated by sequence comparison of the designated clone with the designated GenBank sequence.

Modern: The entries in the "Miden" column of the table refer to the percentage of identically matched with the designated of identically matched in the interpretation of the sequences which is aligned by nucleotides (or residues) that exist along the length of that portion of the sequences which is aligned by

the BLAST comparison to generate the statistical scores presented $\frac{\infty \text{ cvrg}}{(\infty \text{ coverage} = (\text{match} \text{ length} \times 100))}$.

MCBIgi" column.

Table 5. Toxins and Toxin Homologs

NCBI gi description	% Evrg	% Ident	-Assala dorA	Score	jas Gan	NCBI gi	ID NO Gene Id
alpha-latroinsectotoxin precursor - black	EI	67	1.30E-09	135	Score Score	28419952	73 Bt1G100
widow spider (fragment) [Latrodectus							
tredecimguttatus]		•			Ÿ	2.02030	
COMBONENT PRECURSOR HEMOLYSIN BL BINDING	67.	89	1.00E-22	592	330	710702 <u>2</u> g	212 BHC260
(ENTEROTOXIN 40 KD SUBUNIT)					•		
[Bacillus cereus]							
COMBONENT BEETIESOB HEWOLYSIN BL BINDING	8 <i>L</i>	89	1.10E-103	<i>L</i> Z01	066	7107022g	316 Bt1G386
(ENTEROTOXIN 40 KD SUBUNIT) COMPONENT PRECURSOR							
[Bacillus cereus]	•				, , ,		
COMPONENT PRECURSOR HEMOLYSIN BL BINDING	£Ϊ	09	1.10E-10	991	126	7107022g	410 Bt1C203
(ENTEROTOXIN 40 KD SUBUNIT)							

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JERCHARD STEER STE	9	IS	9.60E-05	501	96	ह्हा <u>१</u> ६३	5026 BHC4462
COMPONENT PRECURSOR (D17312) disarbeal toxin [Bacillus cereus] [Bacillus cereus]		15 56	4.70E-169	700 19 44	510 1623	0272681g 7107022g	1222 Bt1G3148
EBacillus cereus) COMPONENT PRECURSOR HEMOLYSIN BL BINDING ENTEROTOXIN 40 KD SUBUNIT)	86	86	7.20E-178	7571	1876	L107022g	419 BilG597
NCBI gi descripti n	ردرو «	% Ident	BlastP- don¶	Score	taa qan Score	NCBI gi	ID NO ZEÓ Ceue Iq

<u>SEQID NO:</u> A sequential SEQ ID NO: is assigned to each contig or singleton and the SEQ ID NO:

corresponds to that set forth in the sequence listing.

Gene ID: Refers to an arbitrarily assigned Gene ID number.

| MCBI gi: Each sequence in the GenBank public database is arbitrarily assigned a unique MCBI gi | Mational Center for Biotechnology Information GenBank Identifier) number. In this table, the MCBI gi number which is associated (in the same row) with a given contig or singleton refers to the particular number which is associated (in the same row) with a given contig or singleton refers to the particular

GenBank sequence which is the best match for that sequence.

sat nap score: The aat nap score is reported by the nap program in the aat package. It is an alignment score in which each match and mismatch is scored based on the BI OSI IM62 scoring matrix.

score in which each match and mismatch is scored based on the BLOSUM62 scoring matrix. Blastp-Prob. Column refer to the probability that such matches occur by

chance. BlastP Score: Each entry in the "BlastP Score" column of the table refers to the BLASTP score that is generated by sequence comparison of the designated clone with the designated GenBank sequence.

generated by sequence comparison of the designated clone with the designated GenBank sequence.

Note that in the "Niden" column of the table refer to the percentage of identically matched nucleotides (or residues) that exist along the length of that portion of the sequences which is aligned by the sequence of the seq

the BLAST comparison to generate the statistical scores presented % cvrg: The %coverage is the percent of hit sequence length that matches to the query sequence

(%coverage = (match length / hit total length) x 100).

**MCBI gi description of the MCBIgi referenced in the MCBI gi description of the MCBIgi referenced in the match in the

"NCBIgi" column.

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